# The effect of paternal factors on perinatal and paediatric outcomes: a systematic review and meta-analysis 

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[^0]BACKGROUND: Maternal factors, including increasing childbearing age and various life-style factors, are associated with poorer shortand long-term outcomes for children, whereas knowledge of paternal parameters is limited. Recently, increasing paternal age has been associated with adverse obstetric outcomes, birth defects, autism spectrum disorders and schizophrenia in children.
OBJECTIVE AND RATIONALE: The aim of this systematic review is to describe the influence of paternal factors on adverse shortand long-term child outcomes.
SEARCH METHODS: PubMed, Embase and Cochrane databases up to January 2017 were searched. Paternal factors examined included paternal age and life-style factors such as body mass index (BMI), adiposity and cigarette smoking. The outcome variables assessed were short-term outcomes such as preterm birth, low birth weight, small for gestational age (SGA), stillbirth, birth defects and chromosomal anomalies. Long-term outcome variables included mortality, cancers, psychiatric diseases/disorders and metabolic diseases. The systematic review follows PRISMA guidelines. Relevant meta-analyses were performed.
OUTCOMES: The search included 14371 articles out of which 238 met the inclusion criteria, and 81 were included in quantitative synthesis (meta-analyses). Paternal age and paternal life-style factors have an association with adverse outcome in offspring. This is particularly evident for psychiatric disorders such as autism, autism spectrum disorders and schizophrenia, but an association is also found with stillbirth, any birth defects, orofacial clefts and trisomy 21 . Paternal height, but not BMI, is associated with birth weight in offspring while paternal BMI is associated with BMI, weight and/or body fat in childhood. Paternal smoking is found to be associated with an increase in SGA, birth defects such as congenital heart defects, and orofacial clefts, cancers, brain tumours and acute lymphoblastic leukaemia. These associations are significant although moderate in size, with most pooled estimates between 1.05 and 1.5 , and none exceeding 2.0.
WIDER IMPLICATIONS: Although the increased risks of adverse outcome in offspring associated with paternal factors and identified in this report represent serious health effects, the magnitude of these effects seems modest.

Key words: paternal age / BMI / smoking / perinatal outcome / birth defects / autism / schizophrenia / cancer / mortality

## Introduction

There is evidence that reproductive failures originate during the periconceptional period and that such failures are influenced by the age and the life-style of the partners (Sinclair and Watkins, 2013; Steegers-Theunissen et al., 2013). Maternal factors, such as the woman's childbearing age and various life-style factors, are associated with poorer short- and long-term outcomes in the offspring (Jacobsson et al., 2004; Wennberg et al., 2016) whereas knowledge of the influence of paternal factors is limited (Soubry et al., 2014).

Over the last few decades, the childbearing age of the mother has increased worldwide from the early 20s to the early 30s (National Institute for Health and Welfare. Nordic perinatal statistics, 2014. Helsinki, Finland). This has increased the focus on the influence of maternal age on the short and long-term health of mothers and children (Jacobsson et al., 2004). However, there has been less focus on the age of the male partner, which has been rising in parallel with maternal childbearing age (Khandwala et al., 2017). Recently, paternal age has been associated with a wide range of adverse health effects in offspring, including both autism spectrum disorders (ASDs) and schizophrenia (Reichenberg et al., 2006). The mechanisms explaining these associations remain unclear (Frans et al., 2015). As a man ages, the number of de novo mutations in his sperm increases along with the chance that a child might carry a deleterious mutation leading to possible diseases (Kong et al., 2012). Recently, a novel mechanism has been suggested which may contribute to the association with paternal age, the process known as 'selfish spermatogonial selection' (Goriely and Wilkie, 2012) where rare spermatogonial cells bearing mutations are positively selected leading to their progressive clonal expansion. This process seems to affect all men, especially as they age.

Furthermore, advanced paternal age has been linked to aneuploidy in autosomes and sex-chromosomes (Lowe et al., 200I; Zhu et al., 2005b)
and epigenetic alterations have been proposed as mechanisms by which modifications in gene expression can be transmitted to the offspring (Perrin et al., 2007). In addition, older fathers may represent a nontypical male population, as both higher and lower socio-economic statuses are overrepresented among older fathers (Nilsen et al., 2013).

Whereas, it is well known that maternal smoking, alcohol consumption and high BMI are associated with poorer short- and longterm outcomes for the children, knowledge of the effects of paternal life-style factors is limited. Malnutrition may impair several metabolic pathways (Steegers-Theunissen et al., 2013), and cigarette smoking can cause DNA or chromosomal damage in human germinal cells, including spermatozoa (Zenzes, 2000). Because ejaculated sperm has minimal, if any, repair capacity it is likely that these changes can be transmitted to the offspring.

The aim of this systematic review was to assess the influence of periconceptional paternal factors on adverse short and long-term child outcomes. These include preterm birth (PTB), low birth weight (LBW), small for gestational age (SGA), birth defects, chromosomal anomalies, psychiatric disorders such as schizophrenia and autism disorders, mortality, impaired neurodevelopment and cognitive functions, and cardio-metabolic functions. We have included the following paternal exposure factors: age, BMI, height, and/or weight and cigarette smoking.

## Methods

We searched the PubMed, Cochrane and Embase databases up to January 2017. Exposures were periconceptional paternal age, paternal smoking and paternal BMI, height, and/or weight. Short-term obstetric outcomes we looked for included PTB, birth weight (BW), LBW, SGA, stillbirth and neonatal death (NND). Further significant outcomes were children with birth defects in general, and selected birth defects i.e.
orofacial clefts, gastroschisis, congenital heart defects (CHDs), spina bifida and trisomy 21 . Long-term outcomes included childhood mortality and morbidity e.g. leukaemia and other malignancies, childhood body weight and BMI, cardio-metabolic disorders, autism/ASD, schizophrenia, other psychiatric disorders and impaired cognitive function. Several of these outcomes when appropriate were used for meta-analysis.

## Systematic search for evidence

The terms used in the searches are listed below:
('Paternal Age'[Mesh]) OR ('Paternal Exposure'[Mesh]) AND ('Congenital Abnormalities')[Mesh] OR congenital malformat*[tiab] OR congenital abnormal*[tiab] OR birth defect*[tiab] OR 'Birth Weight'[Mesh] OR birth weight [tiab] OR birth weight[tiab] OR premature birth[tiab] OR premature delivery[tiab] OR 'Perinatal Mortality'[Mesh] OR 'Perinatal Death'[Mesh] OR perinatal outcome*[tiab] OR ‘Stillbirth'[Mesh] OR ‘Live Birth'[Mesh] OR stillbirth[tiab] OR live birth*[tiab] OR outcome[tiab] OR outcomes[tiab] OR gestational age[tiab] OR children[tiab] OR child[tiab] OR 'Autism Spectrum Disorder'[Mesh] OR 'Autistic Disorder'[Mesh] OR 'Schizophrenia'[Mesh] OR autism[tiab] OR autistic[tiab] OR schizophrenia[tiab]) NOT (animals [mh] AND humans[mh])) NOT ('News'[Publication Type] OR ‘Newspaper Article'[Publication Type]).
We also manually searched reference lists of identified articles for additional references. Guidelines for meta-analysis and systematic reviews of observational studies were followed (Stroup et al., 2000). The literature search was performed by two researchers (C.B. and U.B.W.) and one librarian. Screening of abstracts and of full papers for inclusion was done by pairs of reviewers (C.B. and U.B.W., A.P. and A.L., N.O. and L.B.R., V.S.A. and H.L.). Differences of opinion in the team were solved by discussion until consensus was achieved.

## Inclusion and exclusion of studies

Original studies published in English and the Nordic languages were included. In the case of double publication, the latest study was included. Studies with a control group and case series with more than 100 patients were included. Concerning very rare diseases, studies with fewer cases were also included. Studies published only as abstracts and case reports were excluded. Studies dealing with paternal age were excluded if they did not adjust for maternal age. Systematic reviews without metaanalyses were excluded.

## Definitions

PTB was defined as gestational age $<37$ weeks, very PTB (VPTB) as a gestational age $<32$ weeks. LBW was defined as BW $<2500 \mathrm{~g}$ and very LBW (VLBW) as a BW <1500 g. SGA/intrauterine growth retardation (IUGR), stillbirth and birth defects were defined by each author.

## Appraisal of certainty of evidence

The methodological quality of the studies, in terms of risk of bias, was assessed by pairs of reviewers. We used the tools developed by SBU (Swedish Agency for Health Technology Assessment and Assessment of Social Services) (wnw.sbu.se/sv/Evidensbaserad-vard/Utvardering-av-metoder-i-halso-och-sjukvarden-En-handbok/) for assessing original articles, which grade articles as being of low, moderate and high quality. For systematic reviews we used AMSTAR (AMSTAR: Assessing the Methodological Quality of Systematic Reviews Systematic reviews, cohort and case control studies, but not case series, were assessed for methodological quality. For certainty of evidence we used the Grading of Recommendations Assessment, Development and Evaluation (GRADE) system (Guyatt et al., 2008).

The GRADE system evaluates the following variables for all studies, both combined and per outcome: Design, study limitations, consistency, directness, precision, publication bias, magnitude of effect, relative effect and absolute effect. Certainty levels are divided into high, moderate, low and very low certainty. Certainty levels are based on our confidence in the effect estimate, which in turn is based on the number of studies, design of studies, consistency of associations between studies, study limitations, directness, precision, publication bias, effect size, and relative and absolute effect.

The certainty levels are; very confident $=$ high certainty, moderately confident $=$ moderate certainty, limited confidence $=$ low certainty and very little confidence $=$ very low certainty. If conclusions are based on RCTs, GRADE starts at high certainty level (level 4) but can be downgraded, while if conclusions are based on observational studies GRADE starts at low certainty level (level 2) but might be upgraded (or downgraded). If conclusions were based on case series, no assessment of GRADE was performed.

## Statistics

Outcomes are given in odds ratio (OR), adjusted odds ratio (AOR), adjusted prevalence odds ratio (APOR), hazard ratio (HR), adjusted hazard ratio (AHR), RR (relative risk), adjusted relative risk, adjusted prevalence ratio (APR), adjusted incidence rate ratio (AIRR) or adjusted mortality rate ratio (AMRR) with $95 \% \mathrm{Cl}$. A few studies used mean standardized BW and standardized regression coefficient (Beta).

Meta-analyses were performed despite significant heterogeneity in reference groups for paternal age and despite the fact that outcomes were given in AOR, AHR or APR. A random effects meta-analysis using the DerSimonian and Laird method, with the estimate of heterogeneity being taken from the Mantel-Haenszel model, was used in the analysis (command metan in Stata 15: StataCorp LLC, TX, USA).

## Results

The search strategy identified a total of 14371 articles, of which 238 were selected for inclusion in the systematic review and 81 for inclusion in quantitative synthesis (meta-analysis) (Fig. I, PRISMA Flow chart)

Among the studies included were 10 meta-analyses, 127 cohort studies, 96 case control studies and 5 case series (Supplementary Tables $\mathrm{SI}-\mathrm{III}$ ). Excluded studies, with reasons for exclusion, are presented in Supplementary Table SIV.

A quality assessment of the cohort and case control studies included is presented in Supplementary Tables SV-VII and for systematic reviews in Supplementary Table SVIII. Of the selected cohort and case control studies, 35 articles were of high quality, 103 were of moderate quality and 85 of low quality. Of the systematic reviews included, nine were of medium quality and one was of low quality.

## Paternal age at childbirth and short-term outcomes for offspring

## Obstetric outcomes

PTB and very PTB. Nine cohort studies, comprising more than 10 million births in total, reported on PTB or VPTB or both (Supplementary Table SI, Table I). Six cohort studies were of high quality and three of medium quality. Three studies (Abel et al., 2002; Zhu et al., 2005a; Astolfi et al., 2006) found a small but significantly increased risk of PTB


Figure I PRISMA flow diagram for a systematic review and meta-analysis on the effect of paternal factors on perinatal and paediatric outcomes.
associated with advanced paternal age. Low paternal age (<20 or <25 years) was associated with a higher risk of PTB in four studies (Abel et al., 2002; Chen et al., 2008; Alio et al., 2012; Astolfi et al., 2006). Our meta-analysis including eight of the studies showed a pooled AOR estimate of 1.02 ( $95 \% \mathrm{Cl}$ I.00-I.05) of PTB in older versus younger (reference groups varied between 20 and 34 years) fathers (the forest plot is shown in Fig. 2). In the Danish study by Zhu et al. (2005a) the risk of VPTB in older fathers was increased, but not in the US study by Basso and Wilcox (2006).

Conclusion: There may be little or no difference in the rate of PTB between older and younger fathers. Low certainty of evidence (GRADE $\oplus \oplus O O$ ).

Low BW and very low BW. Nine cohort studies (three high, three medium and three low quality) comprising almost 6 million births assessed LBW, two of them also VLBW (Abel et al., 2002: Chen et al., 2008) (Supplementary Table SI, Table I).

We performed a meta-analysis with LBW as outcome. We included six studies, and found a pooled estimate of $1.00(95 \% \mathrm{Cl}$ $0.97-$ I.03) for LBW in older versus younger (reference groups varied between 20 and 34 years) fathers (Fig. 3). Low paternal age ( $<20$ or
$<25$ years) was associated with a higher risk of LBW in three studies (Abel et al., 2002; Chen et al., 2008; Alio et al., 2012).
None of the studies assessing the risk of VLBW in older fathers found an increased risk.

Conclusion: There may be little or no difference in the rate of LBW between older and younger fathers. Low certainty of evidence (GRADE $\oplus \oplus O O$ ).

Small for gestational age. Five cohort studies (two high, two medium and one low quality), comprising almost 4 million births, found no association between infants born SGA and increased paternal age (Supplementary Table SI, Table I).

Conclusion: There may be little or no difference in the rate of SGA between older and younger fathers. Low certainty of evidence (GRADE $\oplus \oplus O O$ ).

Stillbirth/neonatal mortality. Four cohort studies, comprising between 23821 and 3610647 births, three of high quality and one of medium quality, reported stillbirths (Supplementary Table SI, Table I. In an analysis of more than 3 million births, Astolfi et al. (2004) (high quality) found in an Italian cohort study a significantly increased risk of

Table I Studies on human paternal age and obstetric outcomes identified in a systematic review of the literature on the effect of paternal factors on perinatal and paediatric outcomes.

| Author, year, country | Study design | Number of pregnancies, births or children | Result |  | Outcomes <br> Comments Adjustments | Study quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (risk estimates) | Reference group/ controls |  |  |
| Original articles $\boldsymbol{n}=1 \mathbf{6}$ |  |  |  |  |  |  |
| Abel et al. (2002), USA | Cohort study | 155903 births | Paternal age $<20$ years: <br> PTB: <br> AOR 1.24 (I.02-I.52) <br> LBW: <br> AOR 1.28 (I.02-I.6I) | Paternal age 2I-25 years | PTB and LBW: <br> Significantly higher risk for low paternal age. No other significances. <br> Adjusted for maternal age, socio-economic status, infant gender and race | High |
| Alio et al. (20\|2), USA | Cohort study | 755334 singletons | LBW, PTB (33-37 weeks), VPTB (<33 weeks): <br> Paternal age $<20,20-24$ years: <br> AORs I. 10 to 1.31 (I.07 to I.4I)* <br> $>45$ years: <br> AORs I.I3 to I. 19 (I. 05 to I.44)* SGA: <br> $<20$ years: <br> AORs 1.18 (I.13-1.24) <br> 20-24 years: I.12 (I.10-1.15) <br> Stillbirth ( $\geq 20$ weeks): <br> 40-45 years: AOR I. 24 <br> (1.04-1.47) <br> $>45$ years: 1.33 (1.02-1.77) <br> *lowest and highest $95 \% \mathrm{Cl}$ | Paternal age 25-29 years | LBW, PTB, VPTB: <br> U-shaped risk with significantly higher risk for low and high paternal age. <br> SGA: <br> Significantly higher risk for younger fathers. <br> Stillbirth: <br> Significantly higher risk for older fathers. Adjusted for maternal age, race, education, marital status, years of birth, maternal complications, prenatal care, smoking and alcohol | High |
| Astolfi et al. (2004), Italy | Cohort study | 3619647 births | Stillbirth: <br> Paternal age $\geq 40$ years, maternal age $<35$ years, high education OR I.I2 (I.00-1.25) <br> Paternal age $\geq 40$ years, maternal age $<35$ years, low education: OR I. 29 (I.17-I.43) | Paternal age $<40$ years, maternal age $<35$ years | Stillbirth: <br> Significantly higher risk for high paternal age with low parental education Stratified for maternal age ( $<35$ and $\geq 35$ years) and parental education | High |
| Astolfi et al. (2006), Italy | Cohort study | 1510893 births | Paternal age $<25$ years: <br> PTB: <br> OR I.I9 (I.I2-I.26) <br> VPTB: $1.36 \text { (1.19-1.56) }$ <br> PTB: $35-39,40-44,45-49 \text { years }$ <br> ORs I.OI to I. 36 (I.08 to I.56)* <br> VPTB: $35-39,40-44,45-49 \text { years }$ <br> ORs I.I6 to I. 72 (I. 06 to 2.36 )* <br> *lowest and highest $95 \% \mathrm{Cl}$ | Paternal age 25-29 years, maternal age 25-29 years | PTB, VPTB: <br> U-shaped risk with significantly higher risk for low and high paternal age | High |


| Basso and Wilcox (2006), Denmark | Cohort study | 2499633 live singletons | VPTB: <br> Paternal age 30-34, 35-39, 40-44, 45-49 years: AORs 0.86 to I.II (0.47 to I.55)* Highest AOR among fathers $\geq 50$ years I.3 (0.6-2.8) among women 20-24 years *lowest and highest 95\% Cl | Paternal age 25-29 years, maternal age 25-29 years | No increase in VPTB by paternal age. Stratified for maternal age (20-24, 25-29, 30-34 years), adjusted for mother's education and smoking | High |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Chen et al. (2008), USA | Cohort study | 2614966 singletons | Paternal age $<20$ years: <br> PTB, VPTB, LBW, SGA, NND, <br> low Apgar score: <br> AORs I. 13 to I. 22 (I.OI to I.49)* <br> *lowest and highest 95\% Cl | Paternal age 20-29 years | PTB, VPTB, LBW, SGA, NND, low Apgar score: <br> Significantly higher risk for young age. For high age no significantly increased age up to $>50$ years. <br> Adjusted for paternal race, maternal age, educational level, smoking, alcohol, prenatal care and infant gender | High |
| Iwayama et al. (20II), Japan | Cohort study | 55 005/73993 infants selected at I month and 17263/73 993 at 12 month healthy baby check-up, 3588 underwent both I and I2 months check-up | BW increased with paternal age for non-firstborn infants ( $P=$ 0.0004 ) and LBW decreased with paternal age for non-firstborn infants ( $P=0.0022$ ) | Paternal age was categorized: <20, 20-29, $30-39,40-49$ and $\geq 50$ years. The younger category group was the reference group. | Birth weight: <br> Also included in childhood morbidity. <br> Adjusted for maternal age | Low |
| Nybo Andersen et al. (2004), Denmark | Cohort study | 23821 pregnancies | Late foetal death ( $\geq 20$ weeks): <br> Paternal age $\geq 50$ years: <br> AHR 3.9 (I.I2-I3.8) <br> (3 events), otherwise NS | Paternal age 25-29 years | Stillbirth: <br> Significantly higher high for older fathers. Adjustment for maternal age, reproductive history and maternal life style during pregnancy. Not included in meta-analysis, due to overlap with Urhoj et al. (20I7a) | Medium |
| Olshan et al. (I995), USA | Cohort study | 254892 singletons | No increase in PTB, LBW and SGA by paternal age | Paternal age 25-29 years | PTB, LBW, SGA: <br> Adjusted for maternal age, race, gravidity, smoking, marital status, education and infant gender | Medium |
| Reichman and Teitler (2006), USA | Cohort study | 4621 singletons | Paternal age $>34$ years: LBW: AOR I.7 (I.3-2.2) | Paternal age 20-34 years | LBW: <br> Significantly increased for older fathers. Adjusted for maternal age, gender, mother's birth place, parity, marital status and health insurance status. | Medium/low |
| Selvin and Garfinkel (1972), USA | Cohort study | I 515433 singletons | U-shaped relation with slightly higher rates of LBW at young and older paternal ages | No | LBW: <br> Adjusted for maternal age | Low |
| Stern et al. (20\|4), USA | Cohort study | 9092 ART singletons <br> 6238 subfertile singletons <br> 318816 fertile singletons | Fertile group: <br> Paternal age 35-40, 4I-45, $\geq 46$ years: <br> No association with paternal age and PTB, LBW or SGA | Paternal age $\leq 34$ years, maternal age $\leq 34$ years | PTB, LBW, SGA: <br> Adjusted for parental race and ethnicity, parental education, diabetes, chronic hypertension. Stratified for maternal age, $\leq 34$ and $35-40$ years. | Medium |

## Table I Continued

| Author, year, country | Study design | Number of pregnancies, births or children | Result |  | Outcomes <br> Comments <br> Adjustments | Study quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (risk estimates) | Reference group/ controls |  |  |
| Tough et al. (2003), Canada | Cohort study | 283956 births | Paternal age 20-24, 25-29, 30-34. 35-39, 40-44 years: <br> Significantly decreased risk of LBW: AORs 0.76 to 0.84 ( 0.67 to 0.95)* <br> PTB: AORs 0.75 to 0.87 (0.66 to 0.97)* <br> * lowest and highest 95\% CI | Paternal age $\leq 19$ years | LBW, PTB: Reference group inadequate. Adjusted for maternal age. | Medium |
| Urhoj et al. (2017a), Denmark | Cohort study | 944031 pregnancies with gestational age $\geq 22$ weeks, 4946 stillbirths | Paternal age 35-39, 40-44 and $\geq 50$ years: <br> Stillbirth: AHRs 1.16 to 1.58 (I. 07 to 2.1I)* <br> * lowest and highest $95 \% \mathrm{Cl}$ | Paternal age 30-34 years | Stillbirth: <br> Paternal age associated with the risk of stillbirth in a J-shaped manner with the highest adjusted HRs among fathers $>50$ years. <br> Adjusted for maternal age in I year categories, year of birth: 1994-1999, 2000-2005, 2006-20I0, parental education, in sensitivity analysis also for ethnicity, maternal reproductive history | High |
| Zakar et al. (2015), Pakistan | Cohort study | 5724 births | Paternal age ( $15-24$ years or $\geq 40$ years) was not associated with 'small size at birth' (SSB) or NND | Paternal age 25-39 years | LBW: <br> LBW was a composite of several variables and defined as 'small size at birth (SSB)' (no standard definitions). Controlled for maternal and pregnancy related factors. | Low |
| Zhu et al. (2005a), Denmark | Cohort study | 70347 singletons | PTB: <br> 35-39 years: AOR I.I (I.0-1.3) <br> 40-44 years: AOR I.2 (I.0-1.4) <br> VPTB: <br> 35-39 years: AOR I.4 (I.0-2.0) <br> 40-44 years: AOR I.7 (I.I-2.6) | Paternal age 20-24 years | PTB, VPTB: <br> Risk of PTB, mainly VPTB increased with paternal age <br> Adjusted maternal age, parity, paternal education and income, calendar year and infant gender | High |

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Figure 2 Forest plot describing the association between paternal age and risk for PTB. AOR, adjusted odds ratios.
stillbirth when fathers were 40 or more, compared with fathers below 40 years of age. Alio et al. (2012) also found a significantly increased risk of stillbirth (more than 700,000 US births) for fathers more than 40 years old, when compared to fathers between 25 and 29 years after adjustment for multiple confounders. Two Danish
studies with partial overlap (Nybo Andersen et al., 2004 ~24000 births and Urhoj et al., 2017a almost I million births) found significantly increased risks for the offspring of fathers of 50 years or more (Nybo Andersen HR 3.9, reference group 25-29 years and Urhoj HR I.58, reference group 30-34 years) (Nybo Andersen et al., 2004;


Figure 3 Forest plot describing the association between paternal age and risk for LBW in offspring.

Urhoj et al., 2017a). Our meta-analysis, including three studies, showed a higher risk of stillbirth for the children of older than younger fathers (reference groups varied between 20 and 40 years) with a pooled estimate of I.I9 ( $95 \% \mathrm{Cl}$ I.I0-I.30) (Fig. 4). Two cohort studies reported on NND; one of high quality (Chen et al., 2008) and one of low quality (Zakar et al., 2015). No increased risk of NND was found.

Conclusion: The risk of stillbirth may be slightly increased for older fathers. Low certainty of evidence ( $G R A D E \oplus \oplus O O$ ). It is uncertain whether there is an association between paternal age and NND. Very low certainty of evidence (GRADE円OOO).

## Birth defects and chromosomal anomalies

Children with birth defects. Five studies assessed children with birth defects, three of high quality, one of medium and one of low quality
(Supplementary Table SI, Table II). In a meta-analysis, four of these studies could be included. A small, but significantly higher risk of birth defects was associated with increasing paternal age (pooled estimate I.05, $95 \% \mathrm{Cl}$ I.02-I.07) (Fig. 5). The increase was found already at age 35 years and above.

Conclusion: Higher paternal age is probably associated with a small increase in birth defects. Moderate certainty of evidence (GRADE $\oplus \oplus \oplus O$ ).

Congenital heart defects. Seven studies investigated the association between paternal age and CHDs (Supplementary Table SI, Table II). Five of these studies were high quality, one medium and one of low quality. All studies could be included in the meta-analysis. No significant association was identified between paternal age and CHD (pooled estimate I.03, 95\% Cl 0.99-I.06) (Fig. 6).


Figure 4 Forest plot describing the association between paternal age and risk for stillbirth in offspring.

Conclusion: Higher paternal age is probably associated with little or no difference in the risk of CHD. Moderate certainty of evidence (GRADE $\oplus \oplus \oplus O$ ).

Orofacial clefts. We identified 13 studies, eight of high quality, three of medium quality and two of low quality, for an assessment of orofacial clefts (Supplementary Table SI, Table II). In addition, a systematic review/meta-analysis has analysed the influence of parental age on oral clefts (Herkrath et al., 2012). For paternal age, only two studies could be included in their meta-analysis. A paternal age of 40 years and above was associated with an increased risk of cleft palate (OR $1.58,95 \% \mathrm{Cl}$ 1.15-2.17). We performed a meta-analysis including five studies (Fig. 7). No overall effect of increased paternal age on the incidence of orofacial clefts was found (pooled estimate $0.99,95 \% \mathrm{Cl} 0.95-\mathrm{I} .04$ ) while an age of above 45 years was associated with a small but significant increase in orofacial clefts (pooled estimate I.14, 95\% CI I.02-1.29 (Fig. 7).

Conclusion: Paternal age above 45 years may be associated with a small increase in orofacial clefts. Low certainty of evidence (GRADE $\oplus \oplus O O$ ).

Gastroschisis. Five studies assessed the risk of gastroschisis in association with paternal age, four of high and one of medium quality
(Supplementary Table SI, Table II). All these five studies are included in the meta-analysis (Fig. 8). Overall, higher paternal age was not associated with the risk of gastroschisis (pooled estimate $0.8895 \% \mathrm{Cl}$ $0.78-1.00$ ) (Fig. 8). A significantly lower rate was observed in children whose fathers' paternal ages were between 35 and 40 years, compared to a reference age of between 25 and 29 years ( $U$-shaped association).

Conclusion: Children of fathers aged 35 to 40 years probably have a lower risk of gastroschisis than children of younger fathers. Moderate certainty of evidence (GRADE $\oplus \oplus \oplus$ ).

Spina bifida. Five studies assessed the risk of spina bifida in association with paternal age, three of high and two of medium quality (Supplementary Table SI, Table II). All these five studies could be included in a meta-analysis. No overall higher risk of spina bifida was associated with increasing age (pooled estimate $0.97,95 \% \mathrm{Cl}$ 0.90-1.04) (Fig. 9).

Conclusion: Higher paternal age is probably associated with little or no difference in the risk of spina bifida. Moderate certainty of evidence (GRADE $\oplus \oplus \oplus O$ ).

Chromosomal anomalies. Twenty-three studies of chromosomal anomalies were identified, most of them assessing trisomy 21

## Table II Studies on the association of paternal age with birth defects and chromosomal anomalies in offspring

| Author, year, country | Study design | Number of deliveries or children | Result |  | Outcomes <br> Comments <br> Adjustments | Study quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (risk estimates) | Reference group/ controls |  |  |
| Systematic review |  |  |  |  |  |  |
| Herkrath et al. (2012), Brazil | Systematic review and meta-analysis | 80 articles included in $S R$, 13 articles in metaanalysis, 2 articles about increased paternal age | Paternal age $>40$ years: <br> Cleft palate: OR I.58 (1.15-2.17) | Paternal age 20-39 years | Non-syndromic oral cleft lip with or without cleft palate and cleft palate. Meta-analysis included Harville et al. (2007) and Poletta et al. (2007) <br> Do not specify if adjusted for maternal age | Medium |
| Original articles $\boldsymbol{n}=47$ |  |  |  |  |  |  |
| Archer et al. (2007), USA | Cohort study | Texas Birth Defect <br> Registry. 1996-2002, <br> denominator all live births but number of total births NA <br> Gastroschisis, $n=550$ <br> Trisomy $13, n=143$ <br> Anencephaly, $n=248$ <br> Encephalocele, $n=118$ <br> Trisomy 21, $n=1870$ <br> Cleft palate, $n=2577$ <br> Spina bifida, $n=564$ | Paternal age: <br> Gastroschisis: <br> 20-24 years: APR 1.47 (1.12-1.94) <br> Trisomy 21: <br> 20-24 years: APR I.28 (1.08-1.5। <br> >40 years: APR I. 05 (0.88-I.24) <br> Trisomy I3: <br> $>40$ years: APR 0.40 (0.16-0.96) <br> Cleft palate: <br> 20-24 years: APR I.I8 (1.00-1.38) <br> $>40$ years: APR 0.91 (0.7I-I.16) <br> Pyloric stenosis <br> $>40$ years:APR 0.84 (0.72-0.98) <br> Anencephaly, spina bifida, encephalocele, ASD, VSD (NS) | Paternal age 25-29 years | Selected birth defects. <br> Adjusted for maternal age, race/ethnicity, parity <br> Totally I8.6\% had missing paternal age | High |
| Berg et al. (2015), <br> Norway | Cohort study | 2890 cleft lip (with or without cleft palate) from 2449218 births | Paternal age: <br> 30-34 years: ARR 0.89 (0.80-0.98) <br> 35-39 years: ARR 0.91 (0.79-1.04) <br> 40-44: ARR 0.97 (0.80-1.17) <br> 45-49: ARR I.2I (0.92-I.58) <br> $>50$ : ARR I. 18 (0.78-I.79) | Paternal age 25-29 years <br> Baseline risk I.15/I000 | Cleft lip with or without cleft palate Adjusted for maternal age Interaction analysis showed that the risk was increased only if the age was increased in both parents | High |
| Bille et al. (2005), Denmark | Cohort study | I 489014 births with 1920 non-syndromic (fewer than 3 associated minor anomalies) cleft lip with or without cleft palate +956 cleft palate only | Paternal age 20-50 years: per 10 years increase in age AOR I.I2 (I.02-I.22) for cleft lip with or without cleft palate and AOR I. 24 (I.10-1.40) for cleft palate only | No reference group | Cleft lip with or without cleft palate <br> Adjustment for maternal age Both maternal and paternal ages were associated with the risk of cleft lip with or without cleft palate. For cleft palate alone, only paternal age was a risk factor | High |
| Bunin et al. (1997), USA | Case control | 89 cases | Paternal age (mean $\pm$ SE) <br> $29.9 \pm 0.6$ <br> Age difference 1.5 years, $P=0.07$ <br> versus controls <br> 30-34 years OR $1.8(P=0.05)$ <br> 35-39 years: OR $0.9(P=0.82)$ <br> $\geq 40$ years: OR $2.9(P=0.07)$ | $\begin{aligned} & 178 \text { controls (mean } \pm \mathrm{SE}) \\ & 28.3 \pm 0.5 \end{aligned}$ <br> Reference group: Paternal age $<30$ years | Sporadic neurofibromatosis. Two kind of analyses, one with a control group and another with a reference group Adjustment for maternal age or socioeconomic status did not change the results | Low |
| Cross and Hook (I987), USA | Cohort study | 35680 foetuses with prenatal cytogenetic | No statistically significant effect of any paternal age for trisomy 21 | None | Trisomy 21 <br> Adjustment for maternal age | High |


| de Michelena et al. (I993), Peru | Case control | 318 children and teen agers with trisomy 21 | Means of paternal age for all ages, and for maternal age groups <2I, 2I-29. 30-34, 35-39 and $>39$ years were similar $(P \geq 0.1)$ | 1196 controls (4 controls/ cases) | Trisomy 21 <br> Controls were matched on date of birth, sex and maternal age | Low |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| De Souza et al. (2009), UK | Case control | 471 cases | No statistically significant association between paternal age and trisomy 21, AOR I.I3 (0.85-I.52) per 10 year increase | 456 controls (parents of children with other disabilities) | Trisomy 21 <br> Controls were matched on maternal age (within 3.5 years) and year of birth (within 3 years) | Low |
| De Souza and Morris (2010), UK | Case control | 374 cases with trisomy 13, 929 with trisomy 18, 295 with Klinefelter and 28 with XYY syndrome | Per 10 year increase in paternal age (adjusted for the association of trisomy <br> 21 with paternal age $=$ AOR I. 11 [I.0I-I.23]): <br> Relative (adjusted) to the population <br> Trisomy I3: AOR I.IO (0.83-I.45) <br> Trisomy 18: AOR I.I5 (0.96-1.38) <br> Klinefelter: AOR 1.36 (I.02-I .79) <br> XYY: AOR I. 99 (0.75-5.26) | Population and 5627 controls with trisomy 21 | Trisomy 13, trisomy 18, Klinefelter (XXY), XYY syndrome Controls were matched on maternal age (within 6 months) | Low |
| Dzurova and Pikhart (2005), USA and Czech Republic | Cohort study | Trisomy 21/all births: 593/516745 (California) and 25I/475 834 (Czech Republic) | AOR 0.54-1.02 and NS in California, and AOR 1.49-2.03 and NS in Czech Republic | Paternal age <19 years | Trisomy 21 <br> Adjusted for maternal age, education of mother and sex of infants, 2 years categories Paternal age missing in 17\% in Czech Republic | Medium |
| Erickson and Cohen (I974), USA | Case control | 44/56 cases analysed | Mean paternal age in cases with Apert syndrome: 34.8 years 'Increased paternal age' | Mean paternal age 'population' 32.4 years | Apert syndrome, no statistics given | Low |
| Erickson (1978), USA | Case control | 4000 white infants with trisomy 21 | No independent effect of paternal age (maternal age and birth order constant), rates at paternal age $>45$ years were constant | ‘Some’ 86000 normal white infants | Trisomy 21 | Medium |
| Erickson (1979), USA | Cohort study | 2 data sources, Atlanta data: 226 cases and National Centre for Health Statistics (NCHS) data: I858 cases | Atlanta and NCHS data: no independent paternal age effect using cut off for paternal age $\geq 40,45$ and 50 years | Atlanta data 161452 white and 71 193 black controls, NHCS 4597305 controls | Trisomy 21 | Medium |
| Erickson and Bjerkedal (I98I), Norway | Cohort study | 693 cases | Small age effect in paternal age $\geq 50$ years | 685000 controls | Trisomy 21 <br> Stratified for maternal age <br> No statistics given | Medium |
| Finley et al. (1990), USA | Case control | 14 cases of sporadic blepharophimosis, ptosis, epicanthus inversus, telecanthus complex (BPEI), control data from national means from US statistics | Mean maternal and paternal age higher in cases | US national means 1966-1975 | Blepharophimosis, ptosis, epicanthus inversus, telecanthus complex (BPEI), (autosomal dominant disorder) No statistics given | Low |

## Table II Continued

| Author, year, country | Study design | Number of deliveries or children | Result |  | Comments |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (risk estimates) | Reference group/ controls | Adjustments |  |
| Fisch et al. (2003), USA | Case series | 3419/4387 cases | Effect of paternal age only in mothers $>35$ years ( $P=0.0023$ ) and most pronounced in mothers $>40$ years ( $P=0.0004$ ) | None | Trisomy 21 | Low |
| Green et al. (2010), USA | Case control | Cases with birth defects (if $n \geq 100$ ), between 102 to 6629 cases | Paternal age 40 years <br> All orofacial clefts: AOR I. 07 <br> (0.94-1.23) <br> Septal defects: AOR 0.98 (0.87-I.I0) <br> Spina bifida: AOR I. 03 (0.82-1.28) <br> Omphalocele: AOR 0.92 (0.64-I.33) <br> Gastroschisis: AOR 0.80 (0.54-1.2I) | Control group of 5839 normal live born infants. Paternal age 30 years. Maternal age 28 years | Selected birth defects <br> Adjusted for paternal race and ethnicity, paternal education, maternal alcohol, maternal smoking, parity, earlier miscarriage, plurality, paternal drug used during pregnancy, use of ART, maternal BMI, folic acid use | Medium |
| Grewal et al. (20\|2), USA | Case control | 46 I 14 cases with birth defects | Paternal age: <br> Nervous system anomalies: <br> 38 years: AOR I. 05 (I.00-1.II) <br> 42 years: AOR I.IO (I.02-I.I8) <br> Limb anomalies: <br> 38 years: AOR I. 06 (I.02-I.II) <br> 42 years: AOR I.II (I.05-I.I8) <br> Integument anomalies: <br> 38 years: AOR 1.05 (1.00-1.09) <br> 42 years: AOR I.IO (I.03-I.I6) <br> For fathers 29 years versus $<29$ years: <br> Amniotic band syndrome: AOR 0.87 <br> (0.78-0.97) <br> Pyloric stenosis: AOR 0.93 (0.90-0.96) <br> Anomalies of the great veins: AOR $0.93 \text { (0.87-1.00) }$ | Paternal age <br> 29 year <br> A random sample of 36838 non-malformed births | Selected birth defects Adjusted for maternal age | Medium |
| Harville et al. (2007), Norway | Cohort study | 1431 cases | Paternal age: <br> Cleft palate alone: <br> 30-34 years: AOR I.00 (0.76-1.3I) <br> 35-39 years: AOR 0.94 (0.68-I.30) <br> $\geq 40$ years: AOR I.IO 80.76-I.60) | Paternal age 20-24 years I. 8 million controls | Cleft palate alone Stratified for maternal age, 20-29 years | High |
| Hook et al. (198I), USA | Cohort study | $\begin{aligned} & 55 \text { I cases 1952-1963 and } \\ & 492 \text { cases 1964-1976 } \end{aligned}$ | For 1952-1963 there was no significant paternal age effect (36.87 versus 36.82 years) <br> For 1964-1976, paternal age was about half a year greater in cases than in controls ( 34.55 versus 34.09 years, $P<0.05$ ) | 418017 births 1952-1963 418848 births 1964-1976 | Trisomy 21 <br> Controlled for maternal age | Medium |
| Hook and Cross (I982), USA | Case control | 98 cases of prenatally detected trisomy 21 | Mean difference in paternal age 0.27 $(-1.59 \text { to }+1.06)$ | 10239 foetuses with normal karyotype | Trisomy 21 Controlled for maternal age | Medium |


| Hook and Regal (1984), USA | Case control | 2354 cases with trisomy <br> 21, I 16 cases with trisomy 13 (including cases prenatally diagnosed) | No effect of paternal age | Controls were from all live births the same year in New York State | Trisomy 21 and trisomy 13 associated with Robertsonian translocations Controlled for maternal age and year of birth | Low |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Kazaura and Lie (2002), Norway | Cohort study | \| 738852 infants <br> 1788 trisomy 21 | No increase by paternal age AOR I.II (0.99-I.22) per 10 year increase in paternal age | None | Trisomy 21 <br> Adjusted for maternal age as a continuous variable, birth calendar year and place of birth | High |
| Kazaura et al. (2004a), Norway | Cohort study | 29\| cases with gastroschisis | Higher risk at young paternal age AOR I.6 (I.0-2.4) per 10 year decrease in paternal age after adjustment for maternal age, but was not significant after adjustment for paternal year of birth, AOR per year of father's age: $\text { I. } 04 \text { (0.99-I.08) }$ | >1.7 million controls | Gastroschisis <br> Adjusted for maternal age, paternal year of birth | High |
| Kazaura et al. (2004b), Norway | Cohort study | I 869388 births, 42813 infants with birth defects | Paternal age 45-49 years: <br> CNS defects (not neural tube defects, anencephaly, spina bifida or hydrocephaly) <br> AOR 2.5 (I.2-5.5) <br> Paternal age 20-24 years: <br> Anencephaly AOR I. 4 (I.I-I.8) <br> Neural tube defects. AOR I. 3 <br> (I.I-I.5) <br> Cleft lip: AOR between 0.9-I. 2 (NS) <br> Any birth defects: AOR I.0-I.I <br> (borderline significant) | Paternal age 25-29 years | Any and selected birth defects Adjusted for maternal age, parity, maternity institution and year of birth | High |
| Lian et al. (I986), USA | Cohort study | 7490 infants with a major or serious birth defect | Paternal age $\geq 35$ years versus $<35$ years: <br> ASD: AOR 1.95 (significant) Paternal age $\geq 40$ years versus $<40$ years: <br> Any birth defect: AOR I. 20 (significant) <br> VSD: AOR 1.69 (significant) <br> Chondrodystrophy: AOR I3.32 <br> (significant) <br> Situs inversus: AOR 19.27 (significant) <br> Paternal age $\geq 45$ versus $<45$ years <br> Cleft palate/lip: AOR 2.86 (significant) <br> Trisomy 21: NS any age | 333624 live born control infants without defects | Any and selected birth defects (86 groups of defects) <br> Adjusted for maternal age and race AOR given but no $95 \% \mathrm{Cl}$. | Medium |
| Lorda-Sanchez et al. (I998), Spain | Case control | 14 cases | Associated with increase in: <br> Paternal age ( $34.5 \pm 6.0$ versus $29.6 \pm$ 6.0), OR I.II (I.02-I.2I) but NS after adjustment for maternal age and no of pregnancies | 162 controls | Klippel-Trenaunay-Weber syndrome | Low |

## Table II Continued

| Author, year, | Study design | Number of deliveries | Result |  | Outcomes | Study quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (risk estimates) | Reference group/ controls | Adjustments |  |
| Materna-Kiryluk et al. (2009), Polen | Cohort study | 8683 infants $0-2$ years with birth defects | AOR (per five years increase in paternal age) <br> Heart defects: AOR I. 05 (I.00-I.09) <br> Cleft lip with or without cleft palate: <br> AOR I.II (I.02-I.20) <br> Hypospadia: AOR I.II (I.03-I.19) <br> Gastroschisis: AOR 0.69 (0.54-0.90) | 902452 population | Selected birth defects <br> Adjusted for maternal age | High |
| McIntosh et al. (I995), USA | Case control | 943I cases with 22 different birth defects (Trisomy 2I, $n=997$, cleft palate, $n=1489$ ) | Neural tube defects <br> Paternal age 40-44, $>50$ years AORs 1.6 and 2.3 (borderline significant) <br> Reduction of upper limbs 35-39, <br> 40-44 years <br> AORs: 2.1 and 2.4 (significant) <br> Trisomy 21: <br> 40-44, 45-49, $\geq 50$ years <br> AORs 1.5 to 2.0 (significant) <br> Cleft palate: AORs 0.8-1.5 (NS) | 18862 controls <br> Paternal age 25-29 years | Selected birth defects <br> Adjusted for maternal age | Medium |
| Olshan et al. (1994), USA | Case control | 4110 cases with CHD | General increase by paternal age with trend analysis, NS per age group | 8220 controls <br> Paternal age 25-29 years | CHDs <br> Adjusted for maternal age, no of stillbirths, race | Low |
| Orioli et al. (1995), Italy | Case control | 78 cases with achondroplasia (AC) <br> 64 cases with thanatophoric dysplasia (TD) 106 cases with osteogenesis imperfecta (OI) | Paternal age $>35$ years <br> AC: AOR 3.7I (I.7-8.08) (also <br> mother's age < 30 sign) <br> TD: AOR 3.37 (I.43-8.0) <br> OI: AOR I. 44 (0.78-2.64) | 2 controls per cases <br> Paternal age $<30$ years <br> Maternal age $>30$ years | Achondroplasia (AC), thanatophoric dysplasia (TD), osteogenesis imperfecta (OI), <br> Stratified for maternal age > and <30 years | Low |
| Polednak (1976), USA | Cohort study | 897 orofacial clefts | Total malformation rate: NS <br> Syndactyly: $P<0.05$ <br> Oral cleft rate $1.13 / 1000, P<0.01$ | 776642 population Maternal age 25-29 years | Any and selected birth defects. Stratified for maternal age | Low |
| Poletta et al. (2007), <br> South America | Case control | 5128 cleft lip/palate, 1745 cleft palate | Among 3/II strata (representing 50\% of the cases) significant higher risk with paternal age, ORs between I.42-3.56 | 3712 controls | Orofacial clefts, probably not adjusted for maternal age | Medium/low |
| Riccardi et al. (I984), USA | Case series | 187 cases | Paternal age >35 years: <br> 2 -fold increase | Year matched population controls (from general population). <br> Paternal age $<35$ | Neurofibromatosis Controlled for maternal age | Low |


| Roecker and Huether (1983), USA | Cohort study | 1244 cases | No paternal age effect | 1672210 controls | Trisomy 21 <br> Stratified for maternal age | Low |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Roth et al. (1983a), France | Case control | 118 cases | No effect of paternal age Mean difference in paternal age: 0.46 ( -0.84 to +1.76 ) | 6656 prenatal diagnoses (amniocentesis) | Trisomy 21 Controlled for maternal age | Low |
| Roth et al. (I983b), France | Case series Case control | 2 studies: <br> l:6\| | cases with trisomy <br> 21 <br> 2:242 cases with trisomy <br> 21 | No effect of paternal age Mean difference in paternal age: 0.46 $(-0.34$ to +1.26$)$ | 2 controls per cases in study 2 | Trisomy 21 <br> Controlled for maternal age and time and place of birth | Low |
| Stene et al. (1977), Denmark | Case control | 224 cases | Increased risk of trisomy 21 with paternal age $>55$ years | 5619 controls | Trisomy 21 Controlled for maternal age No statistics | Low |
| Stene et al. (198।), Denmark | Case control | 117 cases | Increased risk of trisomy 21 by paternal age $>4$ I years | 5014 prenatal diagnoses | Trisomy 21 Controlled for maternal age No statistics | Low |
| Su et al. (2015), China | Cohort study (Denmark) | 15216 cases with CHDs | No overall effect of paternal age for CHD <br> PDA (>45 years): AHR I. 69 $(1.17-2.43)$ | I 893899 population Paternal age 25-29 years | CHDs <br> Controlled for maternal age, family history of CHD, maternal infection, gender, parity, parental age difference | High |
| Takano et al. (I992), Japan | Cohort study | 26 cases | No significant effect of paternal age $(P=0.08)$ | Population controls | Neurofibromatosis | Low |
| Tay et al. (1982), Singapore | Case control | 100 cases | No effect of paternal age | 100 controls | Congenital heart disease | Low |
| Tellier et al. (I996), France | Cohort study | 4I cases with CHARGE | Significant higher mean paternal age versus control population <br> Mean paternal age: $33.7+-8$ versus $30.8+-5$ years $(P<0.05)$ | Control population not described | CHARGE malformations (coloboma, heart malformation, choanal atresia, retarded growth, genital hypoplasia, ear anomalies and deafness etc) <br> No difference in maternal age | Low |
| Urhoj et al. (20I5), Denmark | Cohort study | 10817 cases with musculoskeletal congenital abnormalities | AOR I. 06 (I.0I-I.II) by 10 years increase for any musculoskeletal congenital abnormalities $26 \%$ increase ( $2-56 \%$ ) for paternal age $>50$ versus $30-34$ years | I 605885 population | Musculoskeletal congenital abnormalities Adjusted for maternal age, year of birth, ethnicity and education | High |
| Vashist et al. (201I), India | Case series | 200 cases with trisomy 21 | Association with paternal age (correlation coeff $(r)=0.04$, maternal age constant) | Mean paternal age 31.5 years | Trisomy 21 | Low |
| Wolf (1963), USA | Case control | 4II cases with cleft lip and palate | A significant paternal age effect $(P<$ $0.05)$ | 411 controls | Cleft lip and palate Controlled for maternal age | Low |

## Table II Continued

| Author, year, country | Study design | Number of deliveries or children | Result |  | Outcomes <br> Comments <br> Adjustments | Study quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (risk estimates) | Reference group/ controls |  |  |
| Yang et al. (2007), Canada | Cohort study | 77514 cases with birth defects (Trisomy, $n=13078$, cleft palate, $n=6049$ ) | Paternal age 35-39, 40-44, 45-49, $\geq 50$ years <br> Any birth defects: <br> AORs: I.04-I.I5 <br> (significant), test for trend $P=0.015$ <br> Trisomy 21: AORs I.19-I. 45 <br> (significant), test for trend $P<0.01$ Cleft palate: AOR 0.89-1. 23 (NS for any age group) | 5213248 population Paternal age 25-29 years | Any and selected birth defects Advanced paternal age was associated with: Any birth defects, heart defects, tracheooesophageal fistula, oesophageal atresia, musculoskeletal/ integumental anomalies, trisomy 21 and other chromosomal anomalies Paternal age $<25$ years were associated with: Spina bifida/ meningocele, microcephalus, omphalocele/gastroschisis Adjusted for maternal age, race, education, marital status, parity, prenatal care, smoking, alcohol consumption in woman | High |
| Zhan et al. (1991), China | Case control | 497 cases with CHDs | Paternal age $<25$ years AOR:2.63 (2.12-3.27) | 6222 controls <br> Paternal age $\geq 25$ years | Congenital heart disease Controlled for maternal age and birth order | Low |
| Zhu et al. (2005b), USA and Denmark | Cohort study | 3910 cases with birth defects (Trisomy 21, $n=46$ Cleft palate/lip, $n=162$ ) | No overall effect for any birth defect <br> Paternal age $>50$ years: <br> Trisomy 21 <br> AHR: 4.50 (I.0-20.39) <br> Paternal age 35-39 years: <br> Cleft palate: AHR I. 48 (I.02-2.15) <br> Multiple syndromes, extremities, increased by age, test for trend $P<$ 0.00I-0.05 | 7I 937 population Paternal age 20-29 years | Any and selected birth defectsAdjusted for maternal age, parity, maternal and paternal income and education, sex of child and year of birth | High |

[^2]

Figure 5 Forest plot describing the association between paternal age and risk for any birth defects in offspring.
(Supplementary Table SI, Table II). Five of these studies were high-quality cohort studies, nine were medium and nine were low quality, mostly case control studies (Supplementary Table SV). Altogether these studies included more than 37000 ( 37513 ) children with trisomy 21 . Six of the studies could be included in a meta-analysis (Fig. 10). The meta-analysis identified a small but significantly increased risk of trisomy 21 associated with paternal age (pooled estimate $1.13,95 \% \mathrm{Cl} 1.05-1.23$ ). The rate was significantly increased at ages 40 years and above (Fig. 10). For other aneuploidies (trisomy I3 and I8), one high quality study could not identify any increase by paternal age.

Conclusion: Higher paternal age is probably associated with a small increase in the incidence of trisomy 21. Moderate certainty of evidence (GRADE $\oplus \oplus \oplus \bigcirc$ ).

## Paternal age at childbirth and long-term outcomes for offspring

## Morbidity and mortality

Twenty-two studies assessed the effect of paternal age on childhood morbidity including childhood cancer (13 studies), diabetes and obesity
(three studies), developmental disturbances (one study) and mortality (five studies). In addition, one meta-analysis concentrated on the risk of leukaemia (Sergentanis et al., 2015).

Childhood cancer. We identified 13 studies, four of high quality, six of medium quality and three of low quality. They examined childhood cancer in general as well as haematological cancer and leukaemia, retinoblastoma, non-Hodgkin lymphoma, Wilm's tumour and brain tumours in particular (Supplementary Table SI, Table III).

Five studies assessed the effect of paternal age on the incidence of retinoblastoma. Only one study adjusted for maternal age (Yip et al., 2006), and did not report an association between paternal age and retinoblastoma.

In addition, in one systematic review/meta-analysis, the effect of paternal age on the risk of childhood leukaemia was assessed. This showed that higher paternal age was associated with an increased risk of childhood acute lymphoblastic leukaemia (ALL) (RR I.05, 95\% CI I.OI-I.10, per year increments: RR I.04, $95 \% \mathrm{CI}$ I.00-I.08) (Sergentanis et al., 2015).


Figure 6 Forest plot describing the association between paternal age and risk for CHDs in offspring,


Figure 7 Forest plot describing the association between paternal age and risk for orofacial clefts in offspring.

Two studies adjusted for maternal age and could be included in a meta-analysis (Fig. II). No overall higher risk of ALL was associated with increasing age (pooled estimate $1.08,95 \% \mathrm{Cl}$ 0.96-1.21).

Conclusion: Higher paternal age is probably associated with little or no difference in the risk of ALL. Moderate certainty of evidence
(GRADE $\oplus \oplus \oplus O$ ). Higher paternal age may be associated with little or no difference in the risk of other childhood cancers. Low certainty of evidence (GRADE $\oplus \oplus O O$ ).

Diabetes mellitus type I and obesity. Two cohort studies concentrated on paternal age and diabetes mellitus (DM) type I, and one study on


Figure 8 Forest plot describing the association between paternal age and risk for gastroschisis in offspring.
overweight and obesity. Two of these studies were of high quality and one of medium quality (Supplementary Table SI, Table III). In both studies concerning DM, adjustments were made for maternal age. One study (Cardwell et al., 2005) reported an association with paternal age, and the other (Stene et al., 200I) found that paternal age was not significantly associated with DM type I. One study associated an increased risk of obesity with paternal age > 50 years (Eriksen et al., 2013).

Conclusion: It is uncertain whether paternal age is associated with an increased risk of DM type I and obesity in offspring. Very low certainty of evidence (GRADE円OOO).

Mortality. Infant mortality during the first year of life and the possible effects of paternal age were studied by Wunsch and Gourbin (2002), while Urhoj et al. (2014) looked into mortality before the age of 5 years. Three studies concentrated on mortality up to 18,39 , and 40 years, respectively (Zhu et al., 2008; Miller et al., 2010; Mok et al., 2017) (Supplementary Table SI, Table III). In all studies, except Wunsch and Gourbin (2002), adjustments were made for maternal age. These four studies also reported an association of mortality with
higher paternal age. It was not possible to do a meta-analysis because of different length of follow-up in the studies.

Conclusion: Higher paternal age may be associated with a small increase in risk of mortality. Low certainty of evidence (GRADE $\oplus \oplus \bigcirc O)$.

## Psychiatric diseases/disorders in offspring of older fathers

 et al., 2017) and 28 original studies assessed the effect of paternal age on autism and ASD (Supplementary Table SI, Table IV). The original articles included 15 cohort studies and 13 case control studies. Five studies were of high quality, 16 of medium quality and seven of low quality. The systematic review/meta-analysis by Hultman et al. (201I) included 12 studies from seven different countries. The pooled estimates for autism and ASD were for offspring of fathers between 40 and 49 years old 1.78 ( $95 \% \mathrm{Cl} 1.52-2.07$ ), and for offspring of fathers $\geq 50$ years old 2.46 ( $95 \% \mathrm{Cl} 2.20-2.76$ ). In the meta-analysis by Wu et al. (20I7) including 27 studies, a significant association between paternal age and the risk of autism in offspring was found (AOR I.55, 95\% CI I.39-I.73). We included 16 studies in our

Figure 9 Forest plot describing the association between paternal age and risk for spina bifida in offspring.
meta-analysis (Fig. I2). All of these studies adjusted for maternal age. It was observed that there was a higher risk of autism/ASD with increasing paternal age (pooled estimate $1.25,95 \% \mathrm{Cl} 1.20-1.30$ ).

Conclusion: Higher paternal age is probably associated with an increase in autism/ASD. Moderate certainty of evidence (GRADE $\oplus \oplus \oplus$ )

Schizophrenia. Three meta-analyses (Wohl and Gorwood, 2007; Torrey et al., 2009; Miller et al., 2010) and 19 original studies assessed the effect of paternal age on the risk of schizophrenia in offspring. The original articles included 10 cohort studies and 9 case control studies. No studies of high quality, 9 of medium quality and 10 of low quality were included (Supplementary Table SI, Table V). The meta-analysis by Miller et al. (2010) included six cohort and six case control studies. In both study designs, a significant increase in the risk of schizophrenia in the offspring of older fathers was found. The relative risk in the oldest
fathers ( $\geq 50$ years) was 1.66 ( $95 \% \mathrm{Cl} \mathrm{I.46-I.89)} .\mathrm{The} \mathrm{meta-analysis} \mathrm{by}$ Torrey et al. (2009) included ten studies and found an increased risk of schizophrenia in the offspring of the older fathers. The pooled estimate of risk of schizophrenia in offspring of fathers $\geq 55$ years of age was 2.21 ( $95 \% \mathrm{Cl}$ I.46-3.37) and for fathers $\geq 45$ years the pooled estimate was I. 38 ( $95 \% \mathrm{Cl} 0.95-2.0 \mathrm{I}$ ). Wohl and Gorwood (2007) reported an association with paternal age, with higher levels of schizophrenia in the offspring of fathers younger than 20 and older than 35 years.

Fourteen of the original articles were included in a meta-analysis (Fig. 13). All of these studies adjusted for maternal age. Paternal age was categorized as $<35,35-39,40-45,>45$ and $>50$ years. A higher risk of schizophrenia was associated with increasing paternal age (pooled estimate I.3I, 95\% CI I.23-I.38).

Conclusion: Higher paternal age is probably associated with an increased risk of schizophrenia in offspring. Moderate certainty of evidence (GRADE $\oplus \oplus \oplus O$ ).


Figure 10 Forest plot describing the association between paternal age and risk for trisomy 21 in offspring.

Other psychiatric disorders. Fifteen studies concentrated on other psychiatric conditions including attention deficit hyperactivity syndrome (ADHD) (three studies), eating disorders (two studies), psychosis (three studies), bipolar disorders (six studies), Tourette disorder (one
study) and neurocognitive development (one study) (Supplementary Table SI, Table VI).

ADHD Two studies (D’Onofrio et al., 2014; Hvolgaard Mikkelsen et al., 2016) were of high and one of medium quality (Chudal et al.,

## Table III Studies on the association of paternal age with childhood morbidity and mortality in offspring.

| Author, year, country | Study design | Number of children | Result |  | Outcomes | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (Risk estimates) | Reference group/ control |  |  |
| Meta-analysis $\boldsymbol{n}=\mathbf{I}$ |  |  |  |  |  |  |
| Sergentanis et al. (20I5), Greece | Systematic review and meta-analysis | Paternal age as categorical variable: <br> 34 case control studies and 4 cohort studies in systematic review As incremental variable: 9 case control studies and 3 cohort studies in systematic review Meta-analysis (incremental analysis): <br> 8 case control studies and 2 cohort studies | ALL: <br> A 5 year increase in paternal age: RR I. 04 (1.00-1.08) <br> 'Oldest versus middle': <br> Increased risk in oldest fathers with RR $1.10 \text { (1.02-1.19) }$ <br> ARR 1.09 (0.90-1.25) <br> 'Youngest versus middle age': increased risk in youngest with <br> RR I. 09 (I.00-I.20) <br> ARR I. 10 (0.8I-I.5I) <br> AML: <br> No significant associations at incremental analysis or 'oldest versus middle' Increased risk in offspring from 'younger fathers versus middle' with pooled RR I. 28 (1.04-1.59) <br> ARR 0.63 (0.33-1.20) | Paternal age: <br> 'Older': <br> $>35$ years or $>40$ years <br> 'Younger': <br> $<25$ years | Many studies in forest plots did not adjust for maternal age (RR), but sub-analyses regarding degree of adjustments were included (ARR) | Medium |
| Original articles $\boldsymbol{n}=\mathbf{2 2}$ |  |  |  |  |  |  |
| Cardwell et al. (2005), UK Northern Ireland | Cohort study | 991 children with DM type । <br> 447663 cohort | Paternal age: <br> >35 years ARR I.52 (1.10-2.09) | Paternal age <25 years | Children born 197I-I986 Diagnosed with DM type <br> 1 at the age of 15 years <br> Adjusted for maternal age, birth order, year of birth | Medium |
| Crump et al. (20\|2), <br> Sweden | Cohort study | 936 cases with nonHodgkin's lymphoma 3571574 population | Adjusted risk estimates not mentioned in text, no association, $\mathrm{P}_{\text {trend }}=0.3 \mathrm{l}$ | Probably paternal age $<20$ years | Children born 1973-2008 followed through 2009 (ages 0-37 years). <br> Adjusted for perinatal and family variables including maternal age | Medium |
| Crump et al. (2015), <br> Sweden | Cohort study | 2809 cases with brain Tumours 3571574 population (25-29år) | Paternal age: <br> $\geq 35$ years AIRR I. 24 (0.82-I.85) | Paternal age 25-29 years | Children born 1973-2008 followed through 2010 (max age 38 years) <br> Adjusted for maternal age, birth year, sex, foetal growth, parental country at birth, family history of brain-tumour, maternal education | Medium |
| DerKinderen et al. (1990), the NL | Cohort study | 361 sporadic retinoblastoma cases Compared to general population (number NA) | Paternal age: <br> 20-24 years RR 0.4 (no confidence interval available) <br> $>50$ years: RR 5 (no confidence interval available) | Paternal age 25-34 years | Children born 1945-1970 <br> No information about adjustments | Low |
| Dockerty et al. (200I), UK | Case control | 10162 cases with childhood cancer 10162 controls | Paternal age: <br> Retinoblastoma: <br> 40-45 years AOR 0.82 ( $95 \% \mathrm{Cl} 0.39-\mathrm{I} .75$ ) <br> $\geq 45$ years AOR 0.73 ( $95 \% \mathrm{Cl} 0.26-2.0 \mathrm{I}$ ) <br> ALL: <br> 40-44 years AOR I. 45 ( $95 \% \mathrm{Cl}$ <br> I. I0-I.92), $\geq 45$ years AOR I. 54 ( $95 \% \mathrm{Cl}$ (.06-2.23) | Paternal age 25-29 years | Children born 1968-198\| <br> Not adjusted for maternal age | Medium |

## Table III Continued

| Author, year, country | Study design | Number of children | Result |  | Outcomes | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (Risk estimates) | Reference group/ control |  |  |
| Eriksen et al. (20\|3), Norway | Cohort study | 346609 cohort <br> 44712 cases with overweight \| 109 | cases with obesity | Paternal age: <br> Overweight: <br> >50 years ARR I.I5 (0.98-I.35) <br> Obesity: <br> $>50$ years ARR? I.55 (1.14-2.10) | Paternal age $<20$ years | Male conscripts at I8-20 years, born 1967-I984 Overweight defined as BMI $25.0-29.9 \mathrm{~kg} / \mathrm{m} 2$ Obesity defined as BMI $>30 \mathrm{~kg} / \mathrm{m} 2$ Adjusted for birth order, birth years, birth season, maternal age, parity of mother, maternal marital status at birth, parental education level | High |
| Heck et al. (20I2), USA, California | Case control | 609 cases with retinoblastoma 20905 I controls | Paternal age: <br> 30-34 years OR 1.44 (0.99-2.10) <br> $\geq 35$ years OR 1.75 (1.20-2.47) | Paternal age 20-29 years | Children diagnosed with retinoblastoma 1988-2007, children up to 5 years <br> Confounding variables in multivariate models were year of birth, paternal age, urban or rural county of residence, maternal race and maternal place of birth. No information about adjustment for maternal age | Medium |
| Iwayama et al. (2011), Japan | Cohort study | 72268 cohort 16762 cohort attending check-up | Developmental delay AOR 2.25 (1.33-3.80) (36 cases) <br> No word uttering AOR 2.29 (1.52-3.44) <br> (60 cases) <br> Lack of eye contact AOR 5.16 (1.92-13.8) <br> (6 cases) <br> Unable to walk with support AOR I.5I (I.2।-I.91) (230 cases) | Paternal age was categorized: <20, 20-29, 30-39, 40-49 and $\geq 50$ years. The younger category group was the reference group | Children attending child health check-up at age 12 months during 1987-2003 examining child growth and developmental delay. <br> (Included in obstetric outcome, Table Ia) <br> Adjusted for maternal age | Low |
| Johnson et al. (2009), USA | Case control | 17672 cases with childhood cancer 57966 controls | Paternal age: <br> Overall cancer AOR I.OI (0.99-I.03) <br> Leukaemia AOR I. 03 (1.00-1.07) | OR related to a 5 -year increase in paternal age | Children 0-14 years diagnosed with cancer 1980-2004 <br> Controls born 1970-2004 <br> Adjusted for maternal age, sex, birth weight, gestational age, birth order, plurality, maternal race, birth year and state | High |
| Larfors et al. (2012), Sweden | Case control | 2660 cases (children) with leukaemia (AML and ALL) 28288 controls | Paternal age: <br> $\geq 35$ years childhood ALL: <br> AHR I. 08 (95\% Cl 0.95-I.23) <br> $\geq 35$ years childhood AML: <br> AHR 0.89 ( $95 \% \mathrm{Cl} 0.65-\mathrm{I} .22$ ) | Paternal age 20-34 years | Born 1932 or later, diagnosed with leukaemia (children and adult leukaemia) during 1962-2008 Adjusted for sex, Down syndrome and chromosomal aberrations, multiple birth, number of siblings, maternal/paternal age | Medium |
| Matsunaga et al. (1990), Japan | Case control | 225 bilateral and 408 unilateral cases with retinoblastoma Respective 225 and 408 controls | No risk estimate (OR, RR or HR) Only observed and expected numbers | No specific control group | Born during 1965-1968 or 1975-1982 Adjusted by the birth of the children. No adjustments for maternal age | Low |
| Maule et al. (2007), Italy | Cohort study | 229 cases with ALL <br> 284 cases with embryonal tumours 633 I55 population | Paternal age: <br> $\geq 40$ years ALL: ARR 0.93 (0.52-I .67) <br> $\geq 40$ years embryonal tumours: 1.50 (0.90-2.48) | 25-29 years | Children aged I-5 years 1980-1997 <br> Adjusted for sex, year of birth, paternal/maternal age | Medium |


| Miller et al. (2010), Finland | Cohort study | 10965 population 318 cases/deaths | Paternal age: <br> All causes mortality: <br> $\geq 45$ years AHR 3.45 ( $1.85-6.45$ ) <br> Natural death: <br> $\geq 45$ years 2.93 ( $1.05-8.18$ ) | Paternal age 25-29 years | Mortality in offspring born in 1966 and followed to age 39 <br> Adjusted for age of the other parent, subject age, parent social class, maternal parity | Medium |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Mok et al. (2017), <br> Denmark | Cohort study | 179368 I population Cases with premature mortality, psychiatric morbidity and criminality | Paternal age: <br> Natural death: $>45$ years AIRR I. 28 (I.02-1.59) | Paternal age 25-29 years | Mortality in children born 1966-1998 and followed to their 15th or 40th birthday or until 201I Article also in Table 2d: other psychiatric disorders Older paternal age not associated with premature mortality after adjustments except for risk of natural death linked with paternal age 45 years and over. Adjusted for other parent's age, offspring age and sex | Medium |
| Stene et al. (2001), <br> Norway | Cohort study | I 382602 population 1824 with DM type I | Paternal age: <br> $>40$ year ARR I.I6 (0.92-I.46) | Paternal age $<25$ years | DM type I in children born 1974-1998, followed for maximum I5 years until 1989-1998 <br> Adjusted for age group year of birth, maternal age, birth order | High |
| Teras et al. (2015), USA | Cohort study | 2532 cases with haematological malignancies 138003 population | Paternal age: <br> $\geq 35$ år AHR 1.20 (I.0I-I.42) <br> With no siblings: <br> $\geq 35$ years AHR 1.63 (1.19-2.23) | Paternal age $<25$ years | Haematological malignancies diagnosed 1992-2009 No clear linear trend in risk by paternal age Adjusted for age of the other parent and sex | High |
| Urhoj et al. (2014), Denmark | Cohort study | 10855 cases <br> \| 57552 | population | Paternal age: <br> 40-45 years AHR 1.10 ( $95 \% \mathrm{Cl}$ I.00-I.2।) <br> $>45$ years AHR I.I6 (95\% CI I.02-I.32)] | Reference group: <br> Paternal age 30-34 years | Mortality before the age of 5 years in children born 1978-2004 <br> Paternal age associated with increased risk of dying in early childhood due to an excess risk of fatal congenital anomalies, malignancies and external causes. <br> Adjusted for maternal age, parity, parental education, year of birth. | High |
| Urhoj et al. (2017b), Denmark | Cohort study | 3492 cases <br> \| 904363 population | Paternal age: <br> ALL: $\geq 45$ years AHR I. 55 (I.02-2.35) <br> Leukaemias overall: $\geq 45$ years AHR 1.58 (I.07-2.32) | Paternal age 30-34 years | Paternal age associated with risk of ALL with $13 \%$ higher HR for every 5 years increase in paternal age. No firm conclusions for other specific cancer types Adjusted for maternal age, child's year of birth, parental educational levels, parental ethnic origin and maternal parity | High |
| Wunsch and Gourbin (2002), Hungary | Cohort study | 490000 population 8300 cases/deaths | Paternal age: <br> 25-34 years: <br> Neonatal mortality: OR 0.83 (0.76-0.89) <br> Post neonatal mortality: OR 0.99 ( $95 \% \mathrm{Cl}$ $0.95-1.16)$ | Paternal age 35-44 years | Live births and infant deaths from 1984-1988: <br> Early neonatal: up to 7 days of life <br> Neonatal: up to 28 days of life <br> Post neonatal: 28 days to I year of life <br> No adjustments described | Low |
| Yip et al. (2006), <br> Sweden | Cohort study | 7844 cases with cancer: <br> Retinoblastoma $n=226$ <br> Leukaemia $n=1234$ <br> All CNS tumours $n=977$ <br> Astrocytoma $n=316$ <br> Wilm's tumour $n=348$ <br> non-Hodgkin's lymphoma $n=218$ | Paternal age $>40$ years: <br> Retinoblastoma: AIRR 0.96 (0.47-I.97) <br> Leukaemia AIRR I.I4 (0.85-I.53) <br> CNS tumours AIRR I. 69 (1.2I-2.35) <br> Astrocytoma AIRR 1.95 (1.10-3.45) <br> Wilm's tumour AIRR I. 53 (0.89-2.65) <br> Non-Hodgkin's lymphoma $>40$ years AIRR $1.09(0.55-2.16)$ | Paternal age $<25$ years | Children (<15 years) born 1961-2000 with childhood cancer. <br> Adjusted for maternal age | High |

Table III Continued

| Author, year, country | Study design | Number of children | Result |  | Outcomes | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (Risk estimates) | Reference group/ control |  |  |
|  |  | Population 4.3 million |  |  |  |  |
| Zhu et al. (2008), <br> Denmark | Cohort study | 108879 population 831 cases/deaths | Paternal age: <br> 45-49 years: infant mortality AMRR I. 77 $\begin{aligned} & \text { (1.28-2.45) } \\ & \geq 50 \text { år: AMRR } 1.59 \text { (1.03-2.46) } \end{aligned}$ | Paternal age 25-29 years | Mortality in singletons born 1980-1966 followed up to 18 years <br> Adjusted for maternal age and parity, parental education and income, parental country of origin and calendar period | Medium |
| Zorlu et al. (2002), Turkey | Case control | 116 cases with ALL 400 controls | Paternal age: $>40$ years <br> ALL AOR 3.30 (I.28-8.5I) | Paternal age $<40$ years | Diagnosed with leukaemia in 1993-1996, aged I-I4 years. <br> Adjusted for patients' age <br> No adjustments for maternal age (no relationship with maternal age and the risk of ALL was found) | Low |

AIRR, adjusted incidence rate ratio; ARR, adjusted rate ratio; AMRR, adjusted mortality rate ratio;
2015). Chudal et al. (2015) reported an association between younger fathers and ADHD (highest risk associated with paternal age 20-24 years). Another study (D'Onofrio et al., 2014) reported an association of ADHD with advanced paternal age, and the third study (Hvolgaard Mikkelsen et al., 2016) found that there was a higher risk of ADHD if both parents were very young.

Eating disorders Two studies reported on eating disorders (one of high and one of low quality) (Racine et al., 2014; Javaras et al., 2017). The high quality study reported an association between advanced paternal age and eating disorders and also an association with anorexia nervosa (Javaras et al., 2017).

Bipolar disorders Six studies have assessed the risk of bipolar disorders in offspring in relation to advanced paternal age, two study of high quality (Brown et al., 2013; D'Onofrio et al., 2014) three of medium quality (Frans et al., 2008; Chudal et al., 2014; Lehrer et al., 2016) and one of low quality (Menezes et al., 2010). Both Brown et al. (2013) and D'Onofrio et al. (2014) showed that advanced paternal age was a risk factor for bipolar disorders in offspring.

Psychosis and psychotic-like symptoms Three studies, including two case control studies of low quality (Gillberg, 1982; Foutz and Mezuk, 2015) and one study (El-Saadi et al., 2004) which reported results from three different countries, assessed the association between paternal age and psychosis and psychotic-like symptoms in offspring. Two of the studies found an association between advanced paternal age and psychosis (El-Saadi et al., 2004; Foutz and Mezuk, 2015).

Conclusion: It is uncertain whether paternal age is associated with an increased risk of other psychiatric conditions. Very low certainty of evidence (GRADE $\oplus \mathrm{OOO}$ )

## Paternal BMI, height and/or weight at childbirth and short-term outcomes for offspring

## Obstetric outcomes

Altogether 13 cohort studies (mostly of medium quality) have evaluated the effect of paternal BMI, height, and/or weight on obstetric outcomes, in most cases on BW of infants (Supplementary Table SII, Table VII). All studies included in the systematic review had adjusted for maternal factors such as maternal height and BMI. In nine studies the influence of paternal height on BW of the children was studied. In all studies the father's height correlated significantly with BW of the offspring. The effects of BMI, and the weight of the father at the time of conception, or at the beginning of the pregnancy, on neonatal BW were less clear. In one study from 2012, paternal BMI correlated significantly with BW of the newborn, and biparietal diameter, head circumference and pectoral diameter in male offspring (Chen et al., 2012). However, three of six studies did not find any association between paternal BMI and BW of the babies (Table III). Four studies evaluated the correlation between paternal weight at conception and child BW. In three of these reports no association was found (Wilcox et al., 1995; To et al., 1998; Nahum and Stanislaw, 2003). Two studies compared paternal and child BW, with conflicting results (Klebanoff et al., 1998; L'Abee et al., 201 I).

Conclusion: Paternal height is probably associated with BW of the offspring. Moderate certainty of evidence (GRADE $\oplus \oplus \oplus \mathrm{O}$ ). There may be little or no association between paternal $\mathrm{BMI} /$ paternal weight and the BW of the offspring. Low quality of evidence (GRADE $\oplus \oplus O O$ ).


Figure II Forest plot describing the association between paternal age and risk for acute lymphoblastic leukaemia in the offspring.

## Paternal BMI, height and/or weight at childbirth and long-term outcomes for offspring

## Obesity

Paternal anthropometric measurements (BMI, height and/or weight) available at the time of the child's birth were studied in association with childhood outcomes in 13 cohort studies (nine medium and four high quality) and in one medium quality case control study (Supplementary Table SII, Table VIII). In two of the studies paternal height and weight were measured (Durmus et al., 2013; Heppe et al., 2013) and in other studies this information was obtained from questionnaires or records. The outcome was BMI, body fat and/or weight in II studies. Paternal anthropometrics at the time of the child's birth were associated with offspring BMI, weight and/or body fat mass in all studies.

Conclusion: High paternal BMI and weight may be associated with a modest increase in BMI, weight and/or body fat mass in offspring. Low certainty of evidence (GRADE $\oplus \oplus O O$ ).

## ASDs and neurodevelopment

Paternal obesity was an independent risk factor for ASD in children in one medium quality study (Suren et al., 2014) (Supplementary Table SII, Table VIII). In the study of Yeung and co-workers (2017) paternal obesity was associated with delays in personal-social functioning, whereas maternal obesity was associated with delays in fine motor development. Daraki and co-workers (2017) did not find any association between paternal obesity and child neurodevelopment at 4 years of age (Supplementary Table SII, Table IV).

Conclusion: It is uncertain whether there is an association between paternal obesity and ASD and neurodevelopment of the child. Very low certainty of evidence (GRADE $\oplus O O O$ ).

## Paternal smoking at childbirth and shortterm outcomes for offspring

## Obstetric outcomes

Preterm birth. Three cohort studies including more than 30000 children found no increased risk of PTB (<37 weeks) in children where fathers smoked (Supplementary Table SIII, Table IX). Two of the studies were adjusted for maternal smoking (Horta et al., 1997; Ko et al., 2014) and in one cohort study analyses were performed on non-smoking mothers (Andriani and Kuo, 2014). We included three studies in a meta-analysis and found a slight but not significant effect of paternal smoking on PTB (pooled estimate I.16, 95\% I.00-I.35) (Fig. 14).

Conclusion: There may be little or no association between paternal smoking and PTB. Low certainty of evidence (GRADE $\oplus \oplus O O)$.

Low BW. Seven studies (six cohort and one case control), comprising more than 60000 children, investigated the association between paternal smoking during preconception/pregnancy and BW (Supplementary Table SIII, Table IX). In four cohort studies, all adjusted for maternal smoking, no increased risk of LBW was observed in pregnancies where fathers smoked (Horta et al., 1997; Andriani and Kuo, 2014; Ko et al., 2014; Inoue et al., 2016).

Two cohort studies (Magnus et al., I984; Martinez et al., 1994) and one case control study (Zhang and Ratcliffe, 1993) explored the association

Table IV Studies on the association of paternal age with autism/ASDs in offspring.

| Author, year, country | Study design | Number of children | Result |  | Outcomes <br> Comment | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcome | Reference group/ control | Adjustments |  |
| Systematic review and meta-analysis $\boldsymbol{n}=2$ |  |  |  |  |  |  |
| Hultman et al. (2011), <br> Sweden | Meta-analysis <br> (For separate cohort study see under original papers below) | II studies (including 12 cohorts) with paternal age as exposure | Pooled estimates: <br> 30-39 years 1.22 (1.05-1.42) <br> 40-49 years 1.78 (1.52-2.07) <br> $\geq 50$ years 2.46 ( $2.20-2.76$ ) | Paternal age $\leq 29$ years | Increased risk of autism with increased paternal age | Medium |
| Wu et al. (2017), China | Systematic review and meta-analysis | 27 studies (6 cohort and 21 case control studies) with paternal age as exposure | Below reference points: <br> AOR 0.81 (0.73-0.89) <br> Above reference point: <br> AOR I. 55 (1.39-I.73) <br> Overall RR for each 10 year increase in paternal age: $1.21 \text { (1.18-1.24) }$ | Reference point: <br> Midpoint paternal age <br> -age not mentioned | Compared to reference point, lower paternal age was associated with reduced risk and increased paternal age was associated with increased risk of autism | Medium |
| Original articles $\boldsymbol{n}=\mathbf{2 8}$ |  |  |  |  |  |  |
| Ben Itzchak et al. (20II), Israel | Cohort study | 529 cases with ASD Israeli Newborn data, total population not available (2004) | Paternal age in a cohort with ASD. No risk estimate. | Paternal age 20-29 years | The percentage of fathers in the older age ( $30-40$ years) was significantly higher in the ASD cohort compared to the Israeli newborn data ( $P$ $<0,0$ I) | Low |
| Bilder et al. (2009), USA | Case control | 132 cases with ASD <br> 13200 controls | Paternal age: <br> OR I. 28 (0.54-3.03) | Paternal age 30-39 years | No effect of paternal age (high or low) Not adjusted for maternal age | Low |
| Buizer-Voskamp et al. (201I), The <br> Netherlands | Case control | 14231 cases <br> (Autism $n=2262$ <br> Schizophrenia $n=2564$ <br> 8284 Major depression $n=8284$ <br> Bipolar disease $n=$ \| 121 <br> 9048 controls (56 924 controls in total) | Paternal age $\geq 40$ years: <br> Autism: AOR I. 23 (I.OI-I.50) | Paternal age 25-29 years | Adjusted for maternal age $<30$ years and $>30$ years, ethnic background and average income of the residential area | Medium |
| Burd et al. (1999a), USA | Case control study | 78 cases of autism 390 controls | A one-year decrease in the age of the father decreased the risk of autism by $6 \%$ compared to the control <br> OR I. 45 (0.90-2.45) | Paternal age $<20$ or $>30,20-30$ years | Matched controls Increasing father's age associated with increased risk of autism No adjusted risk estimate | Low |
| Byars and Boomsma (2016), Denmark | Cohort study | 10703 cases with autism (ASD) <br> 20586 cases with schizophrenia I 656795 population | Paternal age: <br> 26-30 years AHR 0.92 (0.86-0.98) <br> 35-39 years AHR I.IO (I.03-I.18) <br> 40-44 years AHR I.I9 (I.06-I.33) <br> 45-60 years AHR I.2 (1.04-1.49) | Paternal age 31-34 years | Adjusted for other parent's age (see STable I) | Medium |
| Croen et al. (2007), USA | Case control | 593 cases with autism I32 25I population | Paternal age: <br> 35-39 years AOR I.38 (1.04-1.84) <br> $\geq 40$ years AOR I. 52 (1.10-2.10) <br> With each 10 year increase in paternal age <br> ARR I. 34 (1.06-1.69) | Paternal age 25-29 years | Adjusted for maternal age, birth order, gender, date of birth, parental educational level, ethnicity | Medium |


| D'Onofrio et al. (2014), Sweden | Cohort study | 2424 cases with autism <br> Psychosis, bipolar disorder, suicide attempts 900337 population | Paternal age: <br> $>45$ years AHR I.76 (1.36-2.28) | Paternal age 20-24 years | Adjusted for maternal age, sex, year of birth, parental education, history of psychiatric hospitalization | High |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Durkin et al. (2008), USA | Cohort study | I25I cases with autism 253347 population | Paternal age: <br> $\geq 40$ years AOR I. 4 (I.I-I.8) <br> With each 10 year increase in paternal age AOR I. 3 (I.I-I.5) | Paternal age 25-29 years | Adjusted for maternal age, birth order, gender, maternal education, ethnicity, multiple birth, gestational age, BW for gestational age | High |
| Frans et al. (2013), Sweden | Case control | 5936 cases with autism 30923 controls | Paternal age: <br> 45-49 years AOR I.85 (1.47-2.3।) <br> $\geq 50$ years AOR 2.23 (1.59-3.12) <br> With each 10 years RR I. 25 ( $95 \% \mathrm{CI}$ I.I7, 1.31) | Paternal age 20-24 years | Adjusted for maternal age, birth year, gender, family history, educational level, country | Medium |
| Grether et al. (2009), USA | Cohort study | 23311 cases with autism 7550026 population | Paternal age: <br> 50-54 years AOR I.53 (I.32-I.77) <br> 60-64 years AOR 2.05 (1.38-3.05) <br> With each 10 year increase in paternal age (I5-64 years) ARR I. 22 (I.18, I.25) | Paternal age 25-29 years | Adjusted for maternal age, child's sex, birth weight, ethnicity, education, parity, gestational age, delivery method, birth year | High |
| Hultman et al. (201I), <br> Sweden | Cohort study | 883 cases with autism I 075588 population | Paternal age: <br> 30-39 years AOR I.I9 (I.00-I .42) <br> 40-49 years AOR I. 42 (I.07-I.87) <br> $\geq 50$ years AOR 2.21 (1.26-3.88) <br> $\geq 55$ years AOR 4.36 (2.09-9.09) <br> With each 10 year increase in paternal age RR $1.21(1.10,1.34)$ | Paternal age $<29$ years | Adjusted for maternal age, maternal country of birth, birth weight, maternal history of psychiatric illness, paternal country of birth, paternal history of psychiatric illness, BW, being small/large for gestational age, foetal distress, SES, birth order, year of birth of the offspring | Medium |
| Idring et al. (2014), <br> Sweden | Cohort study | 4746 cases with ASD <br> 417303 population | Paternal age: <br> 25-28 years AOR 0.93 (0.90-0.96) <br> 35-39 years AOR I.07 (I.04-I.IO) <br> 55-59 years AOR I. 39 (I.29-I.50) | Paternal age 32 years | Adjusted for maternal age (using generalized additive models - GAMs), birth year, sex, parity, parental psychiatric history, occupational class, family income, maternal region of birth | High |
| King et al. (2009), USA | Cohort study | 18731 cases with autism <br> 4906926 population | Paternal age: <br> Risk varies over time: <br> >40 years: lowest risk in 1992: RR I. 29 <br> (I.03-1.60) to highest in 1995: RR I.7I (1.4I-2.08) | Paternal age $<30$ years | Autism risk is analysed over multiple birth cohorts <br> Not adjusted risk estimates | Medium |
| Lampi et al. (20\|3), <br> Finland | Case control | 4713 cases with ASD I 132 with childhood autism <br> 1785 with Asperger's syndrome 1796 with pervasive development disorder (PDD) 18777 controls | Paternal age: <br> Autism: <br> 40-49 years AOR I.6 (I.I-2.3) <br> Asperger: <br> 40-49 years AOR I.I (0.5-2.2) <br> PDD: <br> 40-49 years AOR I.6 (0.8-3.2) | Paternal age 25-29 years | Adjusted for maternal age, number of previous births, weight for gestational age, intellectual disability, maternal SES, paternal psychiatric history. | Medium |
| Larsson et al. (2005), <br> Denmark | Case control | 698 cases with autism 17450 controls | Paternal age: <br> >39 years ARR I. 36 (0.96-I.93) | Paternal age 25-29 years | Adjusted for perinatal factors, maternal age, parental psychiatric history, socio-economic characteristics | Medium |

Table IV Continued

| Author, year, country | Study design | Number of children | Result |  | Outcomes <br> Comment <br> Adjustments | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcome | Reference group/ control |  |  |
| Lauritsen et al. (2005) <br> Denmark | Cohort study | 818 cases with autism 943664 population | Paternal age: <br> 40-44 years ARR I.6I (I.19-2.18) <br> $\geq 45$ years: ARR I.2I (1.78-1.86) | Paternal age 25-29 years | Adjusted for maternal age, gender, calendar year of diagnosis, paternal history of psychiatric disease, parental country of birth | Medium |
| Lundstrom et al. (20I0), Sweden | Cohort (twin cohort) - <br> Sweden and UK | Sweden: <br> 164 cases with ASD <br> II 122 population UK: <br> 66 cases with ASD <br> 13524 population | Paternal age in Sweden: <br> 45-50 years AOR I. 90 (1.73-4.92) <br> $\geq 51$ years AOR 3.37 (1.02-11.14) <br> Paternal age in UK: <br> 45-49 years AOR I.66 (0.47-5.82) <br> $\geq 5 \mathrm{I}$ years AOR 3.59 (0.37-34.46) | Paternal age 25-34 years | Twin cohort <br> Adjusted for maternal age, zygosity and SES | Medium |
| Maimburg and Vaeth (2006), Denmark | Case control | 473 cases with infantile autism 4730 controls | Paternal age: <br> >35 years AOR 0.9 (0.7-I.4) | Paternal age 25-29 years | Adjusted for maternal age, maternal citizenship, BW, gestational age, Apgar score, birth defect, irregular foetal position | Medium |
| Mamidala et al. (2013), India | Case control | 471 cases with ASD47। controls | Paternal age: <br> $>30$ years AOR I. 05 (0.76-I.46) | Paternal age $<30$ years | Adjusted for maternal age, gender, year of birth | Medium |
| Parner et al. (2012), Denmark | Cohort study (sibling design) | 9556 cases with ASD <br> \| 31| 736 children | Maternal age $<35$ years and paternal age 35-39 years: <br> AHR 1.27 (1.19-1.35) <br> Maternal age $<35$ years and paternal age $\geq 40$ years: <br> AHR I. 44 (1.3I-I.58) | Paternal age $<35$ years | Combinations of parents' ages: for mothers younger than 35 years, the risk of ASD increased with increasing father's age group. Adjusted for gestational age, birth weight, birth order, sex, parental psychiatric history at birth | Medium |
| Quinlan et al. (2015), USA (New York) | Cohort study | I589 cases with ASD 927003 population | Paternal age: <br> $\geq 35$ years AOR I.4 (I.08-1.68) | Paternal age $<25$ years | Adjusted for parity, sex, race and ethnicity, gestational age, maternal metabolic risk factor, SGA. Not precise whether adjusted for maternal age | Medium |
| Reichenberg et al. (2006), Israel | Cohort study | 319 cases with ASD <br> 13227 \| population | Paternal age: <br> 40-49 years AOR 5.75 (2.65-12.46) <br> With each 10 year increase in paternal age RR $2.14(1.44-3.16)$ | Paternal age 15-29 years | Adjusted for year of birth, SES and maternal age | Medium |
| Sandin et al. (2016), <br> Scandinavia <br> (Danmark, Norge, <br> Sweden), Western <br> Australia and Israel | Cohort study | 30902 cases with ASD <br> 5776794 population | Paternal age and risk for ASD: <br> $<20$ years ARR I. 08 (0.92-I.27) <br> 30-39 years ARR 1.05 (1.02-1.08) <br> 40-49 years ARR I. 28 (1.22-1.34) <br> $\geq 50$ years ARR I. 64 (I.66-I.85) | Paternal age 20-29 years | Joint effect of maternal and paternal age with increasing risk for couples with increasing differences in parental age. <br> Adjusted for site (country), sex, birth year, maternal age | High |
| Sasanfar et al. (2010), Iran | Case control study | 179 cases with ASD 161I controls | Paternal age: <br> $\geq 40$ years AOR 2.03 (1.10-3.73) <br> With each 10 year increase in paternal age a $29 \%$ increase in autism risk | Cohort study: <br> Paternal age 25-29 years <br> Case control study: <br> Paternal age 25-29 years | Adjusted for parental education, birth order, sex, consanguinity, urbanism and province. Not precise whether adjusted for maternal age | Low |


| Shelton et al. (2010), USA | Cohort study | 4947935 population I2 159 cases with autism | Paternal age: <br> >40 years AOR I. 36 (1.26-I.47) | Paternal age 25-29 years | Adjusted for maternal age, parents race/ ethnicity, year of birth, insurance type, parental education | Medium |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Tsuchiya et al. (2008), Japan | Case control | 84 cases with highfunctioning ASD 208 controls | Paternal age: <br> 29-32 years: AOR 2.28 (I.02-5.1I) <br> $\geq 33$ years: AOR 3.09 (1.17-8.16) <br> Paternal age as a continuous variable: AOR $2.54 \text { (0.96-6.72) }$ | Paternal age <29 years | Few cases. <br> Adjusted for maternal age, gender and parity | Low |
| van Balkom et al. (2012), Aruba, The Netherlands | Case control | 95 cases with ASD 347 controls | Paternal age: <br> 30-39 years AOR 2.16 (1.15-4.04) <br> 45-49 years AOR 2.67 (1.07-6.68) | Paternal age $<30$ years | Adjusted for maternal age and PTB | Low |
| Zhang et al. (2010), China | Case control | 95 cases with autism 95 controls | Paternal age: <br> >30 years: AOR 2.63 (1.38-5.00) | Paternal age <30 years | Adjusted for paternal age, gender and birth year. No adjustments for maternal age | Low |

between BW and paternal smoking. Martinez et al. (1994) showed that the number of cigarettes smoked by the father was associated with lower BW in children with non-smoking mothers (test for linearity $P<0.03$ ). This was in line with Zhang, where paternal smoking had a modest effect on BW, the mean BW being 30 grams lower in pregnancies where the fathers smoked (Zhang and Ratcliffe, 1993). In the study by Magnus, paternal smoking had no independent effect on BW in the offspring (Magnus et al., 1984). We performed a metaanalysis including four studies. A small but not significant effect of paternal smoking on the incidence of LBW was found (pooled estimate 1.10, 95\% I.00-1.21) (Fig. I5).

Conclusion: There appears to be little or no association between paternal smoking and LBW. Low certainty of evidence (GRADE $\oplus \oplus O O$ ).

SGA/intrauterine growth retardation. Two cohort studies investigated SGA/ IUGR, defined as BW <10th percentile for gestational age and sex (Horta et al., I997; Ko et al., 2014), comprising more than 27000 children in total (Supplementary Table SIII, Table IX). In Horta et al. (1997) the adjusted risk of SGA/IUGR was significantly increased in pregnancies with paternal smoking, AOR 1.33 ( $95 \% \mathrm{Cl}$ I.05-I.68), while in the study by Ko et al. (2014) no significant association was found (AOR 1.12 ( $95 \% \mathrm{Cl} 0.90-\mathrm{l} .40$ ) for SGA/IUGR. Figures were similar for paternal smoking in the first, second and third trimesters (Ko et al., 2014). A meta-analysis including two studies showed a pooled estimate of I.2I (95\% CI I.03-I.44) (Fig. I6).

Conclusion: Paternal smoking may be associated with a small increase in SGA/IUGR. Low certainty of evidence (GRADE $\oplus \oplus O O)$.

Perinatal mortality. One cohort study of medium quality from Lithuania (comprising 29619 births) including 296 perinatal deaths found an increased risk of perinatal death when fathers smoked (AOR $1.72,95 \% \mathrm{Cl}$ was not available) (Gaizauskiene et al., 2007) (Supplementary Table SIII, Table IX). The probability of foetal and NND was 0.009 in the offspring of fathers who smoked, in comparison with 0.005 in the offspring of non-smoking parents.

Conclusion: It is uncertain whether paternal smoking is associated with perinatal death. Very low certainty of evidence (GRADE $\oplus O O O$ ).

## Birth defects

Eight studies (one cohort and seven case control) reported birth defects in relation to paternal smoking before and during pregnancy (Supplementary Table SIII, Table IX). The cohort study included 14685 births for analysis and found no significant association between paternal smoking and children with orofacial clefts, hydrocephalus, ventricular septal defect (VSD) and urethral stenosis (Savitz et al., 1991). There were seven case control studies. These included a total of 1977 cases, where four studies reported CHD ( $n=1112$ ) (Wasserman et al., 1996; Kuciene and Dulskiene, 2010; Cresci et al., 201 I; Deng et al., 2013), two studies reported orofacial clefts $(n=780)$ (Krapels et al., 2006; Figueiredo et al., 2015) and one study reported anorectal defects ( $n=85$ cases) (van Rooij et al., 2010). Wasserman et al. (1996) also reported neural tube defects ( $n=264$ cases) and limb reduction defects ( $n=178$ cases).

The three CHD studies showed a significantly increased risk associated with paternal smoking with AOR ranging from 1.45 to 3.2 (Kuciene and Dulskiene, 2010; Cresci et al., 201I; Deng et al., 2013), while Wasserman et al. (1996) showed no association (AOR 0.93,


Figure I 2 Forest plot describing the association between paternal age and risk for autism/ASDs in the offspring.

## Table V Studies on the association of paternal age with schizophrenia and schizophrenia spectrum disorders in offspring.

| Author, year, country | Study design | Number of deliveries or children | Result |  | Outcomes <br> Comment <br> Adjustments | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  |
|  |  |  | Outcomes (Risk estimates) | Reference group/ Control |  |  |
| Systematic reviews $\boldsymbol{n}=\mathbf{3}$ |  |  |  |  |  |  |
| Miller et al. (201I), <br> Finland | SR (6 cohort and 6 case control studies) and MA | Cohort studies: 14568 <br> cases and 3000729 <br> controls <br> Case control studies; <br> 8733 cases and <br> 1945092 controls | $\begin{aligned} & \text { Paternal age } \\ & >50 \text { years: RR } 1.66(1.46-1.89) \\ & <25 \text { years: RR } 1.08(1.02-1.14) \text { in male offspring. } \end{aligned}$ | Paternal age 25-29 years | Adjusted for maternal age <br> Sub analyses on gender-stratified data | Medium |
| Torrey et al. (2009), USA | MA (10 studies) | 10 studies included | $\begin{aligned} & \text { Paternal age } \\ & >45 \text { years: OR I. } 38(0.95-2.02) \\ & >55 \text { years: OR } 2.22(1.46-3.37) \end{aligned}$ | NA | Matched for city of birth, season of birth and parental history of treatment for mental disorder | Low |
| Wohl and Gorwood (2007), France | MA (8 studies) | 8 out of 10 studies were included | Paternal age <br> $>54$ years: OR I. 02 (2.03-I7.I2) | Paternal age 25-34 years | Adjusted for maternal age <br> Different age categories used as reference | Low |
| Original articles $\boldsymbol{n}=19$ |  |  |  |  |  |  |
| Brown et al. (2002), USA | Cohort study | 12094 individuals | $73 \%$ response rate (146 of 170 ) <br> Risk for each 10 -year increase in paternal age: ARR I. 89 (I.08-3.32), $Z=2.22, P<0.03$ | Paternal age I5-24 years | Adjusted for maternal age <br> Small study size (7I cases) <br> Includes both schizophrenia and other schizophrenia spectrum diseases (SSD) | Low |
| Buizer-Voskamp et al. (20II), The Netherlands | Case control | 14231 cases 56925 controls | $\begin{aligned} & \text { Paternal age } \\ & >35 \text { age: OR I. } 27 \text { (I.05-I.53) } \\ & \text { Matched controls } \end{aligned}$ | Paternal age 25-29 years | Adjusted for maternal age, SES, and ethnic background <br> Separate analyses for male and female offspring | Medium |
| Byars and Boomsma (2016), Denmark | Cohort study | 1787447 children | 7 out of 15 risk ratios increased in the three age-difference groups Estimates not given | NA | 5 categories for schizophrenic disorders U-shaped association | Low |
| Byrne et al. (2003), USA | Case control | $\begin{aligned} & 7704 \text { cases } \\ & 192590 \text { controls } \end{aligned}$ | Paternal age $>50$ years: Sex-specific estimates. Males with fathers $>55$ years: <br> AIRR 2.10 (I.35-3.28). <br> Females with fathers $>55$ years: <br> AIRR: 3.53 (I.82-6.83) | Paternal age 20-24 years | Adjusted for maternal age, parental education, wealth, marital status and family history of psychiatric history | Medium |
| Dalman and Allebeck (2002), Sweden | Case control | $\begin{aligned} & 420 \text { cases } \\ & 857 \text { controls } \end{aligned}$ | Paternal age <br> >45 years: AOR 2.8 (I.3-6.3) | Paternal age 20-24 years | Brief report. Adjusted for maternal age No adjustment for paternal psychiatric illness | Low |
| Ek et al. (2015), Sweden | Cohort study | $\begin{aligned} & 3829 \text { cases } \\ & 2589502 \text { individuals } \end{aligned}$ | Paternal age <br> >45 years: HR 0.93 (0.72-I.2I) <br> 35-39 years: HR 1.37 (1.18-I.58) <br> 40-44 years: HR I.8I (1.44-2.28) | Paternal age 25-29 years | Adjusted for offspring sex and maternal age Small sample size in the oldest group | Low |
| Frans et al. (201 I), <br> Sweden | Cohort study | 120758 individuals | Paternal age <br> >55 years: AOR I. 95 (1.58-2.40) | Paternal age 20-24 years | Adjusted for maternal age, birth year | Medium |
| Lehrer et al. (2016), USA | Case control | 5317 cases <br> 7658 controls | Paternal age <br> $>45$ years: RRR 2.88 (2.65-3.13) | Paternal age 20-24 years | Adjusted for maternal age <br> Self-reporting of paternal age and clinical history | Medium |
| Malaspina et al. (200I), Israel | Cohort | $\begin{aligned} & 658 \text { cases } \\ & 89722 \text { controls } \end{aligned}$ | Paternal age <br> 40-44 years: ARR I.79 (I.25-2.57) <br> 45-49 years: ARR I.89 (1.24-2.88) <br> $>50$ years: ARR 2.60 (1.63-4.15) | Paternal age 20-24 years | Adjusted for maternal age, sex and ethnic group | Low |
| McGrath et al. (2014), Denmark | Cohort | 2894688 people | Paternal age <br> >45 years: IRR I.54 (I.4I-I.69) | Paternal age 25-29 years | The cohort was observed for 42.7 million personyears | Medium |


| Author, year, country | Study design | Number of deliveries or children | Result <br> Outcomes (Risk estimates) | Reference group/ Control | Outcomes <br> Comment <br> Adjustments | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Naserbakht et al. (201 I), Iran | Case control | $\begin{aligned} & 220 \text { cases } \\ & 220 \text { controls } \end{aligned}$ | Birth rank comparisons: $35 \%$ versus $24 \%$ of the cases versus the controls were in the third or upper birth rank ( $P=0.01$ ). Mean age of fathers at birth in cases ( $30 \pm 6.26$ years) versus controls ( $26.45 \pm 5.64$ years; $P=0.000$ I). Paternal age $\geq 32$ years (at birth) in cases versus controls: AOR 3.8 (I. 80 to 4.27) | NA | Paternal age category is not described Matching for sex and maternal age | Low |
| Petersen et al. (201 I), <br> Denmark | Cohort | 2.2 million people | Paternal age: <br> 45-49 years: AIRR I. 39 (I.13-I.70) <br> 50-54 years: AIRR I. 93 (1.49-2.50) <br> $>55$ years: AIRR I.I5 (I.I2-I-20) | Paternal age 25-29 years | Adjusted for maternal age, proband sex, family psychiatric history in father, mother and siblings The risk of schizophrenia increased with increased paternal age of the father's first child | Medium |
| Sipos et al. (2004), <br> Sweden | Cohort | 754330 people | For each 10 year increase in paternal age: AHR I. 47 (I.23-I.76) | Paternal age 21-24 years | Separate analysis according to family history of the disorder. <br> Adjusted for maternal age, BW, GA, parity and plurality | Low |
| Sorensen et al. (2014), <br> Denmark | Cohort | 176454 men | Cox regression to estimate the IRR of SSD IRR: 1.32 (I.10-I.60) per 10 years increase in paternal age | Paternal age 25-29 years | SSD <br> Adjusted for maternal age, IQ, birth order and family history of psychiatric disorders | Medium |
| Torrey et al. (2009), USA | $\begin{aligned} & \text { Cohort + MA } \\ & \text { (IO studies) } \end{aligned}$ | $\begin{aligned} & 168+88 \text { cases } \\ & 25025 \text { controls } \end{aligned}$ | Cohort of 88 cases: <br> Paternal age: <br> >35 years: OR I. 35 (0.88-2.06) <br> $>40$ years: OR I. 33 (0.75-2.37) <br> $>45$ years: OR I. 32 (0.48-3.63) <br> $\geq 55$ years: <br> MA: pooled OR 2.21 (1.46-3.37) | NA | Matched for city of birth, season of birth and parental history of treatment for mental disorder | Low |
| Tsuchiya et al. (2005), Japan | Case control | 99 cases, 381 controls | ```Paternal age 29-3I years: AOR 2.08 (1.12-3.86) >32 years: AOR 3.00 (1.49-6.04) Test for trend P=0.002``` | Paternal age < 25 years | Adjusted for age and gender of the subject, parity, family history and maternal age. | Low |
| Wang et al. (2015), <br> Taiwan | Case series | 1297 cases | Inverted U-shaped association <br> Onset of schizophrenia was lowered by 1.5 years for paternal age $25-29$ years and by 5.5 years for paternal age $>55$ years Test for trend $P=0.04$ | Paternal age 21-24 years | Study of earlier onset among co-affected sib-pairs with the same familial predisposition. <br> Adjusted for maternal age, gender, education in years and parental education | Low |
| Wu et al. (20I2), China | Case control | 351 cases <br> 238 controls | 351 patients with schizophrenia (167 males, 134 females) Paternal age: <br> $<25$ years: OR 0.628 (0.350-I.I27) <br> 30-34 years: OR 2.660 (1.697-4.169) <br> $>35$ years: OR IO.I83 (4.772-2I.729) | Paternal age 25-29 years | Adjusted for participant's sex, age and maternal age | Low |
| Zammit et al. (2003), <br> Sweden | Cohort | $\begin{aligned} & 2362 \text { cases } \\ & 50087 \text { individuals } \end{aligned}$ | For each IO-years increase in paternal age: AOR I.3 (I.0-I.5); P $=0.015$ <br> Paternal age: 55 years or more: <br> AOR: 3.8 (1.3-II.8) | Paternal age I5-24 years | Adjusted for maternal age, drug use, poor social integration and place of upbringing. |  |



Figure 13 Forest plot describing the association between paternal age and risk for schizophrenia/schizoaffective disorders in the offspring.
$95 \% \mathrm{Cl} 0.58-\mathrm{I} .5$ ). Savitz et al. (I991) found no significant association of paternal smoking and VSD (AOR 2.0, 95\% Cl 0.9-4.3).

We included six studies in a meta-analysis and found a positive association between paternal smoking and CHD (pooled estimate l. 75 (95\% Cl I.25-2.44) (Fig. 17).

There was also a positive association between paternal smoking and orofacial clefts in both case control studies (AOR from 1.45 to 1.5) (Krapels et al., 2006; Figueiredo et al., 2015). However, there was no significantly increased risk in the cohort study, with an APOR I.7 (95\% Cl 0.5-6.0) (Savitz et al., I991). Furthermore, there was a

## Table VI Studies on the association of paternal age with other psychiatric disorders in offspring

| Author, year, country | Study design | Number of deliveries or children | Result |  | Outcomes <br> Comment <br> Adjustments | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (Risk estimates) | Adjustments <br> Reference group/ Control |  |  |
| Original articles $\boldsymbol{n}=15$ |  |  |  |  |  |  |
| Brown et al. (20I3), USA | Case control study | 94 cases <br> 746 controls | Paternal age: <br> Per 10 year increment of paternal age: <br> AOR I. 04 (0.75-I.44) | Paternal age 20-29 years | Bipolar disorders <br> Measured as 10 years increase in paternal age. Controls were matched on date of birth, sex and residency Adjusted for maternal age | Low |
| Burd et al. <br> (I999b), USA | Case control study | $\begin{aligned} & 92 \text { cases } \\ & 460 \text { controls } \end{aligned}$ | Paternal age risk for Tourette: $\beta-0.96, \text { OR } 0.909$ | Each additional year of paternal age decreased the risk of Tourette syndrome by $9.1 \%$ | Tourette syndrome Controls matched for sex, year of birth and month of birth | Low |
| Chudal et al. (2014), Finland | Case control study | I861 cases <br> 3643 controls | Paternal age: <br> $>50$ years <br> AOR 2.84 (I.32-6.I2) <br> 30-34 years <br> AOR I. 35 (I.06-I.72) | Paternal age 25-29 years | Bipolar disorders <br> Adjusted for maternal age and additional adjustment for parental psychiatric history, parental educational level and place of birth | Medium |
| Chudal et al. (2015), Finland | Case control study | $\begin{aligned} & 10409 \text { cases } \\ & 39125 \text { controls } \end{aligned}$ | Paternal age: <br> $<20$ years <br> AOR 1.55 (1.\|I-2.18) <br> 20-24 years <br> AOR 2.20 (1.07-1.34) <br> 45-49 years <br> AOR I. 26 (1.01-I.58) <br> $\geq 50$ years <br> AOR I. 08 (0.73-1.58) | Paternal age 25-29 years | ADHD in singleton births during 1991-2005, diagnosed I995-201। <br> ADHD was associated with young fathers Adjusted for maternal age, paternal psychiatric history, maternal SES, maternal smoking during pregnancy, previous birth, birth weight for gestational age | Medium |
| D'Onofrio et al. (20\|4), Sweden | Cohort study | 2861 cases with ADHD 6819 cases with bipolar disorder 2615081 population | Paternal age $>45$ years <br> ADHD: <br> HR I.76 (1.36-2.28) <br> Sibling fixed-effects model: ADHD: <br> HR I3.I3 (6.85-25.16) <br> Bipolar disorders: <br> HR 24.70 (\|2.12-50.3।) <br> Psychosis: <br> HR 2.07 (I.35-3.20) | Paternal age 20-24 years | ADHD, bipolar disorders, psychosis Adjusted for maternal age, sex, year of birth, parental education, history of psychiatric hospitalization. Also sibling-comparison analyses | High |
|  | Cohort study (Denmark) | Denmark: 11672 cases, <br> 2,3 million population | Paternal age: <br> Denmark: | Paternal age 20-24 years | Psychosis <br> Adjusted for maternal age | Low to medium |


| El-Saadi et al. (2004), Sweden and Australia | Case control (Sweden, Australia) | Sweden: 134 cases and 8687 controls <br> Australia: I 19 cases and 14 \| controls | 35-39 years <br> AOR I. 14 (1.05-1.24) <br> 50-54 years: <br> AOR I.33 (1.33-2.53) <br> $\geq 35$ years: <br> Sweden: <br> AOR 2.42 (1.19-4.89) <br> Australia: <br> AOR 0.77 (0.24-2.46) |  |  | depending on study group |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Foutz and Mezuk (2015), USA | Cohort study | 924 cases <br> 9282 population | Paternal age: <br> 30-34 years: <br> AOR 0,63 (0.25-1.60) <br> $\geq 35$ years: <br> AOR 2.12 (1.08-4.16) | Paternal age 25-29 years | Psychotic-like symptoms <br> Adjusted for demographic characteristics, birth order, lifetime history of depression, anxiety and substance use disorders. | Low |
| Frans et al. (2008), Sweden | Case control study | $\begin{aligned} & 13428 \text { cases } \\ & 67140 \text { controls } \end{aligned}$ | Paternal age: 45-49 years OR 1.14 (I.00-I.30) 50-54 years OR 1.21 (I.00-1.48) $>55$ years OR 1.37 (1.02-1.84) | Paternal age 20-24 years | Bipolar disorders <br> Adjustment for maternal age, additional adjustment for family history of psychotic disorders, parity and socio-economic status | Medium |
| Gillberg (1982), Sweden | Cohort study | 155 cases: <br> Mental retardation 4 <br> Psychosis 30 <br> Psychogenic psychosis 2 <br> Hyperkinetic disorders 3 <br> Anorexia nervosa 5 <br> Conduct disorders 38 <br> Emotional disorders 64 <br> Others 8 <br> 82570 population | Paternal age: <br> No risk estimate | NA | Psychiatric clinic attenders aged 3-19 years during 1975 <br> Psychotic children and adolescents tended to have mothers and fathers who were older than average <br> No adjustments | Low |
| Hvolgaard Mikkelsen et al. (2016), Denmark | Cohort study | $\begin{aligned} & 12294 \text { cases } \\ & 943785 \text { singletons } \end{aligned}$ | Paternal age: <br> 3I-35 years <br> AHR 0.9 (0.77-I.05) <br> $\geq 35$ years <br> AHR 0.74 (0.53-I.02) | Paternal age 26-30 years | ADHD <br> Adjusted for smoking, gender, maternal age | High |
| Javaras et al. (2017), Sweden | Cohort study | 2276809 population <br> Anorexia nervosa $=8137$ <br> Any eating disorder $=16405$ | Paternal age: <br> Anorexia nervosa: <br> 20-24 years <br> AOR 0.9 ( $0.80-0.96$ ) <br> $\geq 45$ years <br> AOR I. 32 (1.14-1.53) <br> Any eating disorder: <br> 20-24 years <br> AOR 0.93 (0.87-0.98) <br> $\geq 45$ years <br> AOR I. 26 (I.13-1.40) | Paternal age 25-29 years | Eating disorders (anorexia nervosa and any eating disorder). Adjusted for sex, birth order, maternal age, country of birth, parental highest education level, lifetime psychiatric and criminal history | High |
|  |  | 5317 cases | Paternal age $\geq 45$ years: |  | Bipolar disorders with or without psychosis | Medium |


| Author, year, country | Study design | Number of deliveries or children | Result |  | Outcomes <br> Comment Adjustments | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (Risk estimates) | Adjustments <br> Reference group/ Control |  |  |
| Lehrer et al. (2016), USA | Case control study | 7658 controls | Bipolar disorder with psychotic features: <br> RRR I. 939 (I.4II-2.643) <br> Bipolar disorder without psychotic features: <br> RRR 0.934 (0.536-I.543) | Paternal age 20-24 years | For schizophrenia and schizoaffective disorders see Table 2c |  |
| Menezes et al. (20I0), Sweden | Cohort study | 493 cases <br> 754330 population | Paternal age: <br> 40-44 years <br> HR I. 85 (I.04-3.30) <br> 45-49 years <br> HR I. 06 (0.39-2.83) <br> $>50$ years <br> HR I. 43 (0.43-4.76) | Paternal age 2I-24 years | Bipolar affective disorders (BPAD) <br> Adjusted for maternal age, SES, family history of psychosis and education <br> Risk of BPAD for each 10-years increase in paternal age | Medium |
| Racine et al. (2014), USA | Cohort study | I722 female twins aged 8-I7 years, II cases | Paternal age: <br> No risk estimate. | Paternal age $\geq 40$ years was coded as reference group for $t$-test comparisons in categorical paternal age models | Eating disorders <br> Advanced paternal age increased the risk for eating pathology Controlled for maternal age | Low |
| Saha et al. (2009), Australia | Cohort study | 33437 singletons | Paternal age 50 years: <br> OR for being in the lowest decile for each neurocognitive variable was significantly associated with elevated paternal age for three of the neurocognitive measures | Paternal age 20 years | Neurocognitive development <br> Adjusted for maternal age, offspring sex, mother's race, weeks of gestation, child's age at testing, family and SES | Medium |

Table VII Studies on the association of paternal BMI, height and weight with obstetric outcomes in offspring.

| Author, year, country | Study design | Number of children | Results | Outcomes <br> Comments <br> Adjustments | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Original articles $\boldsymbol{n}=1 \mathbf{3}$ |  |  |  |  |  |
| Cawley et al. (I954), UK | Cohort study | 1028 children | The height of the fathers was divided into six groups (under 60 inches up to 72 inches and over). BW of the child increased with increased height of father ( $6.73,6.91,7.3 \mathrm{I}, 7.35,7.55$, and 7.74 pounds respectively) | Birth weight (BW) <br> Adjusted for maternal height | Medium |
| Chen et al. (20\|2), China | Cohort study | 889 children; 492 boys and 407 girls | Association between paternal BMI and foetal growth of male offspring: BW $(P=0.0$ I 3$)$, biparietal diameter $(P=0.00$ I $)$, head circumference $(P=$ 0.006), abdominal circumference $(P=0.003)$ and pectoral diameter $(P=0.043)$. Paternal BMI was not associated with foetal growth of female offspring. | BW, newborn's body shape and endocrine system Multivariable regression analysis considering maternal BMI, paternal and maternal age, hypertension during pregnancy, maternal glycated serum protein, parity and gestational age as confounding factors | Low |
| Klebanoff <br> et al. (1998), <br> Denmark | Cohort study | Offspring to girls born in Copenhagen $\begin{aligned} & 1959-196 \mid . \\ & n=3 \mid 30 \end{aligned}$ | Paternal BW was associated with infant BW $(P=0.002)$. <br> Association between paternal adult height and infant BW $(P=0.088)$ Association between paternal BMI and infant BW $(P=0.049)$ | BW <br> Adjusted for maternal BW; maternal adult height, weight, hypertension, diabetes, smoking, education, employment status, and location of residence; child's birth order and gender; other paternal characteristics | Medium |
| L'Abee et al. (201I), the Netherlands | Cohort study | 2947 singletons born 2006-2007 | Paternal BMI and paternal BW were not independent predictors for BW of the offspring | BW <br> Adjusted for maternal factors | Medium |
| Lawlor et al. (2007) Australia | Cohort study | 7223 women and their offspring | Paternal pre-pregnancy BMI: borderline significantly positive association with birth weight standardized for sex and gestational age (regression coefficient 0.03) | BW <br> Paternal coefficient adjusted for maternal effect | Medium |
| Magnus et al. (I984), Norway | Cohort study | 3130 families | Association between paternal weight and infant BW ( $P<0.0 \mathrm{I}$ ) and between paternal height and infant BW ( $P<0.05$ ) | BW <br> Socio-economic status, educational attainment and paternal smoking habit had no independent effects on infant BW | Medium |
| Morrison et al. (1991), <br> Australia | Cohort study | 5989 children | Paternal height was significantly associated with BW $(P<0.0007)$ <br> The increase of the BW was up to 152 g with increased height of the father (ranging from 165 cm to 184 cm ). Paternal BMI had no significant effect on the BW of the child | BW <br> Adjusted for maternal BMI | Medium |
| Mutsaerts et al. (20\|4), Australia | Cohort study | 2264 children | Paternal pre-pregnancy BMI had no influence on PTB AOR 0.99 (0.93-I .06) or SGA AOR 0.96 (0.9I-I.0I) <br> In multivariable analysis paternal BMI did not significantly affect the outcomes | Spontaneous PTB, SGA <br> Adjusted for maternal factors | Medium |
| Nahum and Stanislaw (2003), USA | Cohort study | 24I children | Association between paternal height and child BW $(P=0.02)$ <br> The addition in term BW attributable to each unit increase in paternal height was $10 \mathrm{~g} / \mathrm{cm}$ <br> No significant association between paternal weight and child BW | BW <br> Adjusted for maternal and pregnancy-specific factors | Low |
| Pritchard et al. (1983), UK | Cohort study | 5834 children | The SD scores for BW for taller men were constantly higher than for shorter men. The average difference was 0.29 (approx. 115 g ) for firstborn boys at 40 weeks gestation | BW <br> Adjusted for maternal height | High |


| Author, year, country | Study design | Number of children | Results | Outcomes Comments Adjustments | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: |
| To et al. (I998), China | Cohort study | 355 children born at term | Association between paternal height and child BW $(P<0.01)$ <br> No association between paternal weight and child $(P=0.052)$ <br> No association between paternal BMI and child BW $(P=0.329)$ | BW <br> Analysis of variance of BW adjusted for gestation and controlled for maternal height and pre-pregnancy weight | Medium |
| Wilcox et al. (I995), UK | Cohort study | 571 children | Correlation between paternal height and child BW $(P=0.0115)$ No correlation between paternal weight and child BW $(P=0.2050)$ | BW <br> Adjusted for parental smoking, maternal height, paternal weight |  |
| Winikoff and Debrovner (I98I), USA | Cohort study | 259 children | Paternal height was significantly associated with variations in child BW ( $P<$ 0.05) | BW <br> Adjusted for maternal height, paternal weight, maternal prepregnancy weight, weight during pregnancy | Medium |

significant association between paternal smoking and anorectal malformations (AOR I.8, 95\% CI I.I-2.9) (van Rooij et al., 2010). Our meta-analysis, including two studies of paternal smoking and orofacial clefts, showed a positive association, with a pooled estimate of I.5I ( $95 \% \mathrm{Cl}$ I.I6-I.97) (Fig. I8). However, none of the other birth defects we explored showed a significant association with paternal smoking.

Conclusion: Paternal smoking may be associated with a modest increase in CHD and orofacial clefts. Low certainty of evidence (GRADE $\oplus \oplus O O)$.

## Paternal smoking at childbirth and longterm outcomes for offspring

## Cancer

Five studies explored the association between paternal smoking during pregnancy and cancers in offspring (Supplementary Table SIII, Table X ). Of these, three studies divided smoking into two sharply distinguished classifications, smoking/non-smoking, but only reported dose-response estimates (ji et al., I997; Sorahan et al., 200 I; Pang et al., 2003). In the use of cigarettes $>5$ pack-years, Ji et al. (1997) found a significant association between paternal smoking and cancer in offspring (AOR I.7, $95 \% \mathrm{Cl}$ I.2-2.5). Likewise Sorahan et al. (200I) showed a significant association between smoking and cancer: 10 to 19 cigarettes per day (AOR I.63, $95 \% \mathrm{Cl}$ I.IO-2.4I) and 20-29 cigarettes per day (AOR I.46, (95\% CI I.05-2.03). However, Pang et al. (2003) did not show an association between cancer and the father smoking $>20$ cigarettes per day. In the two studies with the smoking/non-smoking dichotomy, Sorahan and Lancashire (2004) showed a significant association between smoking and cancer in offspring (AOR I.28, $95 \% \mathrm{Cl}$ I.I5-I.42), while John et al. (199|) did not. Childhood acute leukaemia and brain tumours are dealt with in the sections below, while paternal smoking was not associated with any of the specific cancers in any of the studies.

Conclusion: Paternal smoking during pregnancy may be associated with a modest increase in cancer in offspring. Low certainty of evidence (GRADE $\oplus \oplus O O$ ).

Acute childhood leukaemia. Out of 19 original studies, two were of medium and 17 of low quality (Supplementary Table SIII, Table $X$ ). Studies rated as low quality included only a few cases, and the information on paternal smoking in the preconception period and during pregnancy was collected retrospectively from mothers several years after birth. The majority of studies found no association between maternal smoking and childhood leukaemia, hence they did not adjust for maternal smoking in the analyses of paternal smoking.

Acute lymphoblastic leukaemia. Two meta-analyses on paternal smoking and ALL have been published (Liu et al., 20II; Milne et al., 20I2). Milne et al. (2012) included both a meta-analysis and original data in their paper (Table X ). All 10 studies included in the meta-analysis by Milne et al. (2012) were also included in the meta-analysis by Liu et al. (2011), except for the original data: the latter meta-analysis included 18 case control studies. Thirteen studies explored paternal smoking during the preconception period (AOR I.25, $95 \% \mathrm{Cl}$ I.08-I.46) and eight studies during pregnancy (AOR I.24, $95 \% \mathrm{Cl}$ I.07-I.43). Their dose-response analysis estimated a higher risk

## Table VIII Studies on the association of paternal BMI, height and weight with long-term outcomes in offspring

| Author, year, country | Study design | Number of children | Results | Outcomes <br> Comments <br> Adjustments | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Original studies $\boldsymbol{n}=14$ |  |  |  |  |  |
| Catalano et al. (2009), USA | Case control study | 89 children (52 from NGT and 37 from GDM pregnancies | Paternal weight at the time birth was greater in children in tertile 3 of weight percentiles compared to children in tertile I at follow-up at $8.8 \pm 1.8$ years No difference in paternal BMI at the time of child's birth in relation to tertiles of percentage body fat of the child | Childhood weight and body fat (measured by dual energy $X$-ray absorptiometry) at follow-up <br> Reference: Centers for Disease Control and Prevention (CDC) weight and body fat percentiles <br> Maternal obstetrical data, paternal anthropometric data and neonatal birth data was included to best determine which combination of perinatal factors best modelled the risk of adiposity in child Maternal pre-pregnancy BMI was the strongest predictor of childhood obesity | Medium |
| Cawley et al. (1954), UK | Cohort study | 1028 children | Infant weight was more highly correlated with height of mother than father. Correlation of infant height and weight at 24 months; mother 0.21 , father 0.13 | Weight of 625 children with observations at all intervals (6,9,12 and 24 months) <br> Adjusted for maternal height | Medium |
| Daraki et al. (2017), <br> Greece | Cohort study | 772 children | Paternal obesity not associated with child neurodevelopment at 4 years of age | Child neurodevelopment <br> Adjusted for maternal BMI | Medium |
| Davey Smith et al. (2007), UK | Cohort study | 4654 children | The association between paternal BMI and offspring BMI at 7.5 years of age:0.202 standardised age and sex adjusted coefficient ( $0.175-0.229$ ), similar to maternal BMI | Childhood BMI <br> Standardised regression coefficients age and sex adjusted Sensitivity analysis for non-paternity performed Maternal and paternal BMI were included in the same model. | High |
| Durmus et al. (201I), The Netherlands | Cohort study | 5674 children | Pre-pregnancy paternal BMI was strongly associated with childhood overweight at the age of 4 years <br> The main effects of maternal BMI on childhood BMI were stronger than the main effects of paternal BMI ( $P<0.00$ I; and $P=0.013$, respectively) <br> As compared to children from parents with normal BMI, children from two obese parents had an increased risk of overweight at the age of years, OR 6.52 (3.44-I2.38) | Childhood height, weight and BMI. <br> Maternal BMI had a significantly stronger effect on childhood BMI Adjusted for maternal BMI | High |
| Heppe et al. (2013), The Netherlands | Cohort study | 3610 children | Higher paternal BMI associated with higher risk of preschool overweight: OR 1.35 (I.19-I.53) | Preschool overweight <br> Values reflect the OR and $95 \% \mathrm{Cl}$ for each parental or child characteristics that remained in the backward selection model. Maternal pre-pregnancy BMI in the model | High |
| Jaaskelainen et al. (201I), <br> Finland | Cohort study <br> (NFBC <br> 1986) | 4788 children | Paternal pre-pregnancy obesity strongly predicted overweight: <br> At 16 years of age: <br> Father-son OR 3.17 (I.70-5.92) <br> Father-daughter OR 5.58 (3.09-10.07) <br> If both parents obese, overweight of the child at 16 years: <br> Sons OR 5.66 (3.12-10.27) <br> Daughters OR 14.84 (7.4I-29.73) | Childhood overweight <br> Long-term overweight of both parents had a greater impact on the risk of offspring overweight than parental weight gain from normal weight to overweight/obesity during the 16 -year follow-up period | High |
| Lawlor et al. (2007), <br> Australia | Cohort study | 7223 children | The increase in standardized offspring BMI at age 14 for a one SD increase in paternal BMI was 0.239 SD (0.197-0.282) | Childhood BMI <br> The maternal-offspring BMI association was stronger than the paternaloffspring BMI association <br> Paternal coefficient adjusted for maternal effect | Medium |

Table VIII Continued

| Author, year, country | Study design | Number of children | Results | Outcomes <br> Comments <br> Adjustments | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Lawlor et al. (2008), UK | Cohort study | 4091 children | As assessed at 9 to 11 years of age, mean difference in offspring sex- and agestandardized fat mass $z$-score per I SD BMI $0.24(0.22-0.26)$ for maternal BMI versus $0.13(0.11-0.15)$ for paternal BMI | Offspring fat and lean mass (measured by dual energy X -ray absorptiometry) <br> Adjusted for maternal BMI <br> Maternal effect size association was larger | Medium |
| Linabery et al. (2013), USA | Cohort study | 912 children | Infants of obese fathers had BMI growth curves distinct from those of normal weight fathers. The $p$ value for the global association between paternal BMI category and infant BMI growth curves (birth- 3.5 years) from joint mixed effect model was 0.02 | Infant BMI <br> Maternal BMI has a stronger influence on BMI growth than paternal BMI Missing exposure (II\% maternal and $26 \%$ paternal BMIs) and covariate data were assumed to be missing at random and imputed, outcomes were not imputed <br> Adjusted for maternal BMI in the joint model | Medium |
| O'Callaghan et al. (1997), Australia | Cohort study | 4062 children | Paternal BMI is independent predictor of severe and moderate obesity at 5 years of age. <br> Paternal BMI percentiles 85-94: <br> Severe obesity RR 2.8 (1.8-4.5) <br> Moderate obesity RR I. 0 (0.6-I.5) <br> Paternal BMI percentiles $>95$ : <br> Severe obesity RR 2.0 (I.1-3.6) <br> Moderate obesity RR 2.1 (1.4-3.3) | Offspring obesity (BMI class) <br> Reference category: paternal BMI percentiles 15-84 | Medium |
| Reilly et al. (2005), UK | Cohort study | 7758 children | Paternal obesity was associated with the risk of obesity in children at 7 years of age. Final model AOR: Father $(\mathrm{BMI}>30) 2.54$ (I.72-3.75) compared to both parents with $\mathrm{BMI}<30 .$ <br> As compared to children from parents with normal BMI, children from two obese parents had an increased risk of overweight at the age of 4 years: OR 6.52 (3.44-12.38) | Offspring obesity based on BMI <br> Maternal and paternal BMI were entered in the logistic regression models with risk of severe or moderate obesity as the dependent variable | Medium |
| Suren et al. (20\|4), Norway | Cohort study | 92909 children | ASD in children at the age of 4.0-13.1 (mean 7.4) years: <br> Paternal BMI $>30$ : versus $\mathrm{BMI}<25$ <br> AOR 1.73 (I.07-2.82) <br> Asperger disorder in children aged $\geq 7$ years: <br> Paternal BMI $>30$ versus $\mathrm{BMI}<25$ <br> AOR 2.01(1.13-3.57) | ASDs <br> Adjusted for maternal BMI | Medium |
| Yeung et al. (20I7), USA | Cohort study | 4821 children | Increased risk of failing the personal-social domain in children up to 3 years of age Paternal BMI > 30 compared with children of normal weight fathers AOR I.7I (I.08-2.70) <br> Children whose parents both had BMI $\geq 35$ were likely to additionally fail the problem-solving domain | Delays in childhood development Adjusted for maternal obesity | Medium |

ASD, autism spectrum disorder; GDM, gestational diabetes mellitus; NGT, normal glucose tolerance; RR, relative risk;

## Table IX Studies on the association of paternal smoking with obstetric outcomes and birth defects in offspring

| Author, year, country | Study design | Number of deliveries or children | Result |  | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (Risk estimates) | Comment <br> Adjustments |  |
| Obstetric outcomes $\boldsymbol{n}=8$ |  |  |  |  |  |
| Andriani and Kuo (2014), Taiwan | Cohort 1993-2007 | 3789 children <br> 71.5\% paternal smoking with non- <br> smoking mothers <br> $2.4 \%$ only mother smoking | LBW <br> Only father smoking during pregnancy AOR 0.89 (0.5I-I.54) <br> I-IO cig/day AOR 0.8 I (0.58-I.14)* <br> II-20 cig/day AOR 0.66 (0.46-0.94)* <br> $\geq 20$ cig/day AOR 2.09 (1.38-3.17)* <br> PTB <br> Only father smoking during pregnancy AOR I. 16 (0.78-I.7I) <br> I-IO cig/day AOR 0.5 I (0.34-0.75)* <br> II-20 cig/day AOR 0.78 ( $0.55-\mathrm{I} .1$ I)* <br> $\geq 20$ cig/day AOR 2.11 (1.38-3.23)* | Questionnaires to both parents Adjusted for sex, birth order, maternal age, father's education, maternal employment status, parental BMI, household income, urban/rural residence *Only adjusted for birth order | Medium |
| Gaizauskiene et al. (2007), Lithuania | Birth Registry study 2002 | 296 Perinatal death <br> (199 stillborn +97 died within the first six days after delivery) <br> Total cohort $N=29619$ | Perinatal death <br> Paternal smoking <br> AOR I.6 (I.4-2.2) <br> Paternal smoking without maternal smoking <br> AOR I. 72 (no Cl) | Confounders 45 parameters <br> Maternal age $<$ or $\geq 36$ years <br> Education, marriage/cohabiting | Low |
| Horta et al. (I997), Brazil | Cohort 1993 | 5166 singleton live birth 2237 (43.3\%) paternal smoking | LBW: AOR 1.18 (0.94-I.48) PTB: AOR I. 25 (0.99-I.57) IUGR: AOR I. 33 (I.05-I.68) | Mothers interviewed soon after delivery by trained interviewers Adjusted for social class, maternal schooling, parity, birth interval, prior LBW, maternal height, number of antenatal care visits and for maternal smoking | Medium |
| Inoue et al. (2016), Japan | Prospective hospital cohort study 1997-2010 | Total 21855 newborn <br> Present study 16396 <br> Non-participants 5459 (25\%) <br> Smoking: <br> 5905 Mother no/Father yes | LBW <br> Smoking only fathers 502 children with LBW <br> AOR I. 07 (0.94-I.22) | Birth after GA 37 weeks <br> Mothers interviewed in I. trimester <br> Adjusted for maternal smoking, paternal smoking (or four types of combination of interaction effect), maternal age, paternal age, maternal BMI, maternal occupational status, parity, sex | Medium |
| Ko et al. (2014), Taiwan | Birth Cohort study 2005-2006 | Total 24200 children Interview rate 87.8\% Included 21 248 children | LBW, PTB, SGA <br> Preconception $>20$ cigarettes per day: <br> PTB, $n=101 / 1158$ (8.7\%) <br> AOR I. 07 (0.84-1.35) <br> LBW, $n=84 / 1158$ (7.3\%) <br> AOR I. 14 (0.87-1.27) <br> SGA, $n=114 / 1 \mid 58$ (9.8\%) <br> AOR I.I2 (0.90-I.40) | Interview 6 months post-partum (mothers) <br> Adjusted for maternal age, nationality, education, parity, total weight gain during pregnancy, gender of infant, multiple birth and maternal smoking in the same period Similar results for smoking in 1. or 2-3 trimester | Medium |
| Magnus et al. (I984), Norway | Cohort study 1967-1979 | 3130 singletons <br> II I75 pairs of like-sexed twins born 1915-1960 <br> Singletons born 1967-1979 | Birth weight <br> Correlation matrix <br> Paternal smoking <br> Regression coefficients (+/-SE) <br> Bivariate regression -48 (8.9) $P<0.0$ I | Paternal height and weight, Maternal height and weight, paternal and maternal education, SES, maternal smoking Correlation matrix | Low |

## Table IX Continued

| Author, year, country | Study design | Number of deliveries or children | Result |  | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Outcomes (Risk estimates) | Comment <br> Adjustments |  |
| Martinez et al. (I994), USA |  |  | Multiple regression -4.9 (9.3) NS |  |  |
|  | Cohort study | 1596 eligible children 350 refused (22\%) <br> 1219 respondents <br> 191 father smoking <br> 992 mother non-smoking <br> 907 non-smoking mothers in multiple regression analyses | Birth weight <br> Paternal with non-smoking mother: <br> None: $3602 \mathrm{~g} \pm 401$ <br> I-10:3573 g $\pm 388$ <br> $11-20: 3520 \mathrm{~g} \pm 382$ <br> $>20: 3514 \mathrm{~g} \pm 428$ <br> $P$ for linearity 0.026 <br> 88 g adjusted BW difference <br> $P$ for linearity $=0.026$ | Hospital data at birth + questionnaire $\leq I$ months after birth by parents <br> Multiple regression analysis for non-smoking Mothers. Adjusted for GA, birth order, ethnicity, maternal and paternal education, maternal age, sex | Medium |
| Zhang and Ratcliffe (I993), Shanghai | Case control 1986-1987 | 1785 full term singleton live born normal infants of non-smoking mothers 1033 exposed to paternal smoking 752 non-exposed | Birth weight <br> 3236 g versus 3262 g <br> Birth weight diff 26 g (NS) <br> Multiple linear regression $30 \mathrm{~g}(95 \% \mathrm{Cl}=-7.66)$ | Pretested in-hospital interview to mothers after delivery Adjusted for parity, maternal age, gestational age, maternal occupation. <br> Modestly adverse effect on birth weight | Low |
| Birth defects $\boldsymbol{n}=8$ |  |  |  |  |  |
| Cresci et al. (20II), Italy | Case control 2008-2010 | $\begin{aligned} & 360 \text { cases } \\ & 360 \text { controls } \end{aligned}$ | CHD <br> Paternal smoking OR I.7(I.I-2.6) <br> Paternal smoking $\geq 15$ cigarettes per day: <br> OR 2.1 (1.3-3.5) | Questionnaires to both parents. Matched for age range Unconditional regression adjusted for potential confounders but not specified <br> Maternal smoking insignificant but not directly adjusted for | Low |
| Deng et al. (20\|3), China | Case control $2010-2011$ | 284 cases <br> 422 controls <br> Subgroups <br> Light smoking I-9 cig/day <br> Medium IO-19 cig/day <br> Heavy $\geq 20$ cig/day | Non-syndromic CHD <br> Paternal smoking during peri-conception period (3 months before pregnancy and first trimester) and non-smoking mothers <br> No avoidance behaviour: <br> Septal defects: 37 cases/46 controls <br> AOR 2.52 (1.39-4.59) <br> Cono-truncal defects: 36 cases/46 controls <br> AOR 3.22 (1.75-5.93) <br> Other outcomes listed but less than 20 cases | Maternal face-to-face interview Adjusted for residence, age, education, pre-pregnant BMI, alcohol use, folic acid use, paternal alcohol and family history of CHD | Low |
| Figueiredo et al. (2015), USA | Prospective case control Democratic Republic of Congo, Vietnam, Philippines, Honduras | 430 cases <br> 754 controls <br> Fathers smoking <br> 173 cases <br> 245 controls | Orofacial cleft <br> Paternal smoking 3 months before and during pregnancy <br> AORI. 5 (I.I-I.9) | Interview of mothers <br> Age $<3$ years <br> Adjusted for sex, parental employment and education and age, location at birth rural/city, country <br> Maternal smoking infrequent - only I.4-I.9\% | Medium |
| Krapels et al. (2006), The Netherlands | Case control | $\begin{aligned} & 350 \text { cases } \\ & 222 \text { controls } \\ & \text { Fathers smoking }>10 \text { cig/day } \\ & 62 \text { Cleft lip }+/- \text { cleft palate } \\ & 17 \text { Cleft palate only } \end{aligned}$ | Orofacial cleft <br> Univariate analyses paternal smoking: <br> $>10$ cigarettes per day: <br> Cleft lip with or without cleft palate: OR I. 5 (1.0-2.4) <br> Cleft palate only: OR I. 8 (0.9-3.5) | Questionnaires to both parents <br> Follow-up two years after the <br> peri-conception period <br> Multivariate analyses are not shown for paternal smoking but was not significant | Low |
|  | Case control register study | 261 newborns | Congenital heart septal defects |  | Low |

Interviews of both parents, mostly mothersAdjusted for maternal education, social status, and marital status Medium 3
Interviews of both parents, mostly mothersAdjusted for
maternal education, social status, and marital status
Interview by mothers at first prenatal care visit
Adjusted for maternal age, race, education and maternal
smoking Questionnaire to parents
Median age: 5-6 years
Multiple adjustments but estimate not confounded by any
covariate
Maternal smoking insignificant but not directly adjusted for.
Telephone interview of mothers
Paternal smoking I month before through 3 months after
conception.
Risk estimates were adjusted for selected non-specified
covariates did not differ substantially from crude estimates. covaniates did ? Only father smoking
AOR I.45(I.03-2.03)
Congenital anomalies
Paternal smoking
Cleft lip with or without cleft palate: POR I.7
(0.5-6.0)
Hydrocephalus: POR 2.4 (0.6-9.3)
Ventricular septal defect: POR 2.0 (0.9-4.3)
Urethral stenosis: POR 2.0 (0.6-6.4)
Anorectal malformations
Paternal smoking 3 months before conception:
OR I.8 (I.I-2.9)
Birth defects
Father only smokers:
Cono-truncal heart: $35 / 90$ OR 0.93 (0.58-I.5)
Neural tube: 59/90 OR I.I (0.76-I.7)
Limb reduction defect: 4 I/90 OR I.4 (0.88-2.2)
associated with an increased number of cigarettes a day (CPD), $>20$ CPD (AOR I.30; 95\% Cl I.09-I.55) (Liu et al., 201 I). Milne et al. (2012) also found a significantly increased risk of ALL when fathers smoked around the time of conception (AOR I.I5, 95\% CI I.06-I.24). For $>20$ CPD their meta-analysis included seven studies (AOR I.44, $95 \% \mathrm{Cl} 1.24-\mathrm{l} .68$ ).

Three low quality studies out of the 17 original studies in this systematic review were not included in the meta-analysis. CastroJimenez and Orozco-Vargas (201I) included 85 matched pairs and found AOR 1.93 ( $95 \% \mathrm{Cl}$ I.06-3.54), Farioli et al. (2014) included only risk estimates according to the following categories $1-10 \mathrm{CPD}$ (AOR 0.86, $95 \% \mathrm{Cl} 0.58-\mathrm{l} .26$ ) and $>10 \mathrm{CPD}$ (AOR 0.74 ( $95 \% \mathrm{Cl}$ $0.5 \mathrm{I}-\mathrm{l} .05$ ). The Metayer et al. (20|3) study was an expansion of Chang's (Chang et al., 2006) (included in both meta-analyses) and found an AOR 0.94 ( $95 \% \mathrm{Cl} 0.69-1.27$ ). Based on the meta-analyses, paternal smoking was associated with a $15-25 \%$ increased risk of ALL.

Conclusion: Paternal smoking may be associated with a slightly higher risk of childhood ALL. Low certainty of evidence (GRADE $\oplus \oplus \bigcirc \bigcirc)$.

Acute myeloid leukaemia. Twelve original studies (two of medium quality and 10 of low quality) evaluated the outcome of paternal smoking on acute myeloid leukaemia (AML) (Supplementary Table SIII, Table X). The meta-analysis included eight studies and two unpublished reports. Figures for paternal smoking prior to conception were AOR I.I9 (95\% $\mathrm{Cl} 1.00-\mathrm{I} .4 \mathrm{I}$ ) and during pregnancy AOR I. 28 ( $95 \% \mathrm{Cl}$ I.05-I.57). All original studies in our systematic review are included in the metaanalysis.

Conclusion: There appears to be little or no association between paternal smoking and childhood AML. Low certainty of evidence (GRADE $\oplus \oplus O O$ ).

Brain tumours. Fourteen studies explored the association between paternal smoking prior to and during pregnancy, and brain tumours (Supplementary Table SIII, Table X). All were included in our metaanalysis, which showed a significant association between paternal smoking and brain tumours (pooled estimate $1.12,95 \% \mathrm{Cl}$ 1.03-I.22) (Fig. I9).

Conclusion: Paternal smoking may be associated with a small increase in childhood brain tumours. Low certainty of evidence (GRADE $\oplus \oplus \bigcirc \bigcirc)$

Cardio-metabolic outcomes. Nine cohort studies of medium quality assessed paternal smoking and cardio-metabolic outcomes in offspring (Supplementary Table SIII, Table X). Five studies examined BMI (Leary et al., 2006; Kwok et al., 2010; Durmus et al., 201I; Howe et al., 2012; Florath et al., 2014), three studies looked at blood pressure and hypertension (Brion et al., 2007; de Jonge et al., 2013; Taal et al., 2013), and one study explored DM type I (Toschke et al., 2007). Due to the heterogeneity of the studies, meta-analyses could not be performed for any of these outcomes.
The BMI of children was measured at various ages in the five studies and the results diverged. In two of the studies, no linear associations were observed between paternal smoking and BMI (Durmus et al., 201I; Howe et al., 2012). However, three studies showed a negative linear association between paternal smoking and BMI in children aged 7 to 10 years (Leary et al., 2006; Kwok et al., 2010; Florath et al., 2014).


Figure 14 Forest plot describing the association between paternal smoking and risk for PTB.

Brion et al. (2007) and Taal et al. (2013) showed no association between paternal smoking and diastolic or systolic BP in offspring in the adjusted models, neither did de Jonge et al. (2013) show any association between paternal smoking and hypertension in daughters. Toschke et al. (2007), in two combined cohorts, showed significantly lower risk estimates of DM type I in children where fathers smoked during pregnancy (AOR $0.44,95 \% \mathrm{Cl} 0.25-0.75$ ).

Conclusion: It is uncertain whether there is any association between paternal smoking and BMI in offspring. Very low certainty of evidence (GRADE $\oplus O O O$ ). There may be little or no association between paternal smoking and blood pressure in offspring. Low certainty of evidence ( $G R A D E \oplus \oplus O O$ ). It is uncertain whether there is any association between paternal smoking and DM type $I$ in the offspring. Very low certainty of evidence (GRADE $\oplus O O O$ ).

## Neuro-developmental outcomes

Six cohort studies explored the association between paternal smoking and neuro-developmental outcomes, out of which three cohort studies studied ADHD (Nomura et al., 2010; Langley et al., 2012; Zhu et al., 2014) (Supplementary Table SIII, Table X). All studies were heterogeneous regarding child age, questionnaires used for the parents, and outcome measures. Two of the studies found a significant association between paternal smoking and ADHD (AOR ranging from 1.29-I.42) (Langley et al., 2012; Zhu et al., 2014), while Nomura et al. (2010) found no significantly increased risk (AOR 0.3I, 95\% Cl 0.06-I.92). In Langley et al.
(2012) and Zhu et al. (2014) the children were 7 and 8 years of age, while the children in Nomura et al. (2010) were only 3 and 4 years old. None of the other studies found any associations between paternal smoking and neuro-developmental outcomes, except Brion et al. (20|0), who found a significant association between paternal smoking and conduct/externalizing problems (AOR I.12, 95\% Cl I.02-I.24).

Conclusion: Paternal smoking may be associated with a small increase in ADHD. Low certainty of evidence (GRADE $\oplus \oplus O$ ). It is uncertain if there is any association between paternal smoking and other neuro-developmental outcomes. Very low certainty of evidence (GRADE $\oplus O O O)$.

## Discussion

## General discussion

In this systematic review and meta-analysis we have tried to summarize the evidence for the effect of paternal factors on perinatal and paediatric outcomes. Paternal factors investigated in the present paper were paternal age and life-style factors, in particular smoking and $\mathrm{BMI} /$ height/weight of the fathers at time of conception. Other exposures, such as male subfertility and teratogenic drugs, have not been included in the present systematic review. Table XI presents a summary of the findings from the meta-analyses.


Figure 15 Forest plot describing the association between paternal smoking and risk for LBW in offspring.


Figure 16 Forest plot describing the association between paternal smoking and risk for SGA in offspring.


Figure I7 Forest plot describing the association between paternal smoking and risk for CHDs in offspring.


Figure 18 Forest plot describing the association between paternal smoking and risk for orofacial clefts.

## Table X Studies on the association of paternal smoking with long-term outcomes in offspring.

| Author, year, country | Study design | Number of deliveries and children | Result <br> Outcomes <br> (Risk estimates) | Outcomes <br> Adjustments | Quality assessment |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Cancer <br> Acute childhood leukaemia Meta-analyses $\boldsymbol{n}=\mathbf{3}$ |  |  |  |  |  |
| Metayer et al. (2016), USA | SR with meta-analysis Pooled analyses of CLIC studies | Childhood Leukaemia International Consortium (CLIC) studies: Meta-analyses including 6-9 CLIC studies and 3-4 Non-CLIC studies Pooled analysis of 12 case control studies with 1330 AML <br> 13169 controls | AML <br> Paternal smoking during preconception period MA (CLIC+non-CLIC): AOR 1.19 (I.00-I.4I) Pooled CLIC studies: AOR I.I8 (I.0I-I.38)* Paternal smoking during pregnancy MA (CLIC+non-CLIC): AOR I. 28 (I.05-1.57) Pooled CLIC studies: AOR I. 24 (1.06-1.46)* Paternal ever smoking <br> MA (CLIC+non-CLCI): AOR I.I8 (0.92-I.5I) Pooled CLIC studies: AORI. 34 (I.II-I.62)* | Interview with mothers and/or fathers, age < 15 yrs <br> Adjusted for age, sex, ethnicity, paternal education, study centre <br> *Similar results for analyses including only non-smoking mothers (data not shown) <br> Dose-response relationship with paternal smoking <br> Maternal smoking had no effect in pooled CLIC analysis or meta-analysis and is not adjusted for <br> High correlation between pre-and postnatal paternal smoking. Limited ability to identify specific windows of exposure | Medium |
| Milne et al. (20I2), Australia | SR and meta-analysis 10 case control studies, including own study | Any versus none: 10 studies 5338 cases <br> Controls: NA >20 CPD:7 studies 2118 cases Controls: NA | ALL <br> Paternal smoking around the time of conception: <br> Any versus none: OR 1.15 (1.06-1.24) >20 CPD: OR 1.44 (I.24-1.68) | All studies except own study by Milne et al. (2012) are also included in the meta-analysis by Liu et al. (201I) | Low |
| Liu et al. (201I), USA | SR and meta-analysis, I8 case control studies | Preconception: <br> I3 studies <br> Cases and controls: NA | ALL <br> Paternal smoking during preconception: <br> AOR I. 25 (I.08-I.46)* <br> Paternal smoking during pregnancy: <br> AOR I. 24 (I.07-I.43) <br> Dose-response a. > 10 CPD; b. 10-19; c. $>20$ <br> a. AOR I.I7 (0.9-I.54) <br> b. AOR 1.25 (I.0I-I.55) <br> c. AOR 1.30 (1.09-I.55) | Primarily interviews by mothers <br> Age 18 month to 18 years <br> Most studies matched and adjusted for potential confounders *Only 5 studies included in MA adjusted for maternal smoking Also, a positive association between ALL and paternal ever smoking and at each exposure time period examined | Medium |
| Original articles $\boldsymbol{n}=19$ |  |  |  |  |  |
| Brondum et al. (I999), USA | Case control (CCG study) 1989-93 | 1618 ALL <br> 1722 controls 450 AML <br> 523 controls | ALL <br> Paternal smoking I month before pregnancy <br> AOR I. 07 (0.90-1.27) <br> Father (not mother) ever smoked ( $n=1842$ ) <br> AOR I. 04 (0.86-I.26) <br> AML <br> Paternal smoking I month before pregnancy AOR 0.87 (0.64-1.18) <br> Father (not mother) ever smoked ( $n=517$ ) <br> AOR I. 32 (0.9।-I.93) | Telephone interview with parents mostly mothers <br> Child age: <br> ALL <15 years, AML <18 years <br> Matched by age, race, telephone code area <br> Adjusted for annual income, father's and mother's exposures, race and education <br> No association with maternal smoking, parental years of smoking, or number of pack-years | Medium |
|  | Case control | 85 cases | ALL | Face-to-face interview with parents | Low |

## Table X Continued

| Author, year, country | Study design | Number of deliveries and children | Result Outcomes (Risk estimates) | Outcomes <br> Adjustments | Quality assessment |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Castro-Jimenez and OrozcoVargas (201I), Colombia | 2000-2005 | 85 controls | Paternal preconception smoking AOR 1.93 (I.06-3.54) | Age $<15$ years <br> Matched control sex, age, region <br> Not adjusted for maternal smoking but of no significance |  |
| Chang et al. (2006), USA | Case control 1995-2002 (Northern California Childhood leukaemia study) | 281 ALL <br> 46 AML <br> 416 controls <br> Paternal preconception <br> smoking <br> ALL: <br> 74 cases <br> 70 controls <br> AML: <br> 16 cases <br> 8 controls | ALL <br> Paternal preconception smoking AOR I. 32 (0.86-2.04) <br> AML <br> Paternal preconception smoking AOR 3.84 (1.04-14.17) | Self-administered questionnaire/in-person interview of mothers <br> Age <15 years <br> Matched on age, maternal race, and Hispanic ethnicity. <br> Adjusting for household income <br> Maternal smoking was not associated with increased risk of ALL or AML <br> Data included in Metayer et al. (2013) | Low |
| Farioli et al. (2014), Italy | Case control 1998-2003 (SETIL study) | 557 cases 855 controls <br> I-I0 CPD: <br> 77 cases <br> 108 controls <br> $>10$ CPD: <br> \| 51 cases <br> 222 controls | ALL <br> Only paternal smoking I-IO CPD in conception period AOR 0.86 (0.58-I.26) $>10$ CPD in conception period AOR 0.74 (0.51-।. 05 ) | Personal interview with parents <br> Age < 10 years <br> Mutually adjusted models <br> also including paternal smoking during pregnancy and maternal smoking in first trimester <br> Child second-hand-smoking (SHS), birth order, BW, duration of breast feeding, mat and pat age, educational level, birth year mother, parental exposure benzene | Low |
| ji et al. (I997), China | $\begin{aligned} & \text { Case control } \\ & \text { \|98\|-\|99\| } \end{aligned}$ | 642 cases <br> 642 controls <br> No maternal smoking <br> Acute leukaemia <br> 166 case control pairs <br> Lymphoma <br> 87 case control pairs | Cancer <br> <2 Pack-years (PY) <br> 2-5 PY <br> >5 PY prior to conception <br> Acute leukaemia AOR 2.4 (I.I-5.6)* <br> ALL AOR 3.8 (1.3-12.3)* <br> AML AOR 2.3 (0.4-14.8)* <br> Lymphoma AOR 4.5 (1.2-16.8)* <br> All cancers AOR I.7 (I.2-2.5)* | Paternal and maternal interviews by trained interviewers Age < 15 years. Matched for sex, year of birth Adjusted for BW, income, paternal age, education and alcohol For $<5$ PY there were no significant risk in any of the cancers | Low |
| John et al. (1991), USA | Case control 1976-83 | 223 cases <br> 196 controls | Cancer <br> Paternal smoking preconception period, <br> absence of maternal smoking <br> ALL: AOR I.4 (0.6-3.I) <br> Lymphomas: AOR I.6 (0.5-5.4) <br> Brain cancer: I. 6 (0.7-3.5) <br> All cancers: AOR I.2 (0.8-2.1) | Personal interview Prenatal exposure Age 0-14 years Matched for age, sex, area Absence of maternal smoking: Adjusted for father's education. | Low |
| Lee et al. (2009), Chorea | $\begin{aligned} & \text { Case control } \\ & \text { 2003-2005 } \end{aligned}$ | 164 cases leukaemia 106 ALL <br> 164 controls | All leukaemia and ALL <br> PY before pregnancy <br> All leukaemia: >IO PY: AOR I.7 (0.9-3.3) <br> ALL: > IO PY: AOR I. 6 (0.8-3.5) | Interview with mothers (93.5\%) <br> Age 0-18 years <br> Matched for age and sex. <br> Adjusted for age, gender, father's education and birth weight Maternal smoking was too small ( $6.1 \%$ in controls) to be evaluated in childhood leukaemia risk and was not considered further | Low |
|  | Case control | 399 cases | Acute leukaemia | Personal interviews with each child parents | Low |


| MacArthur et al. (2008), Canada | (the cross-Canada childhood leukaemia study) 1990-1994 | 399 controls <br> 109 cases <br> 96 controls | Paternal smoking before pregnancy: <br> <10 CPD <br> 10-19 CPD <br> $\geq 20$ CPD <br> All leukaemia: <br> AOR 0.99 (0.50-I.99) <br> AOR I.I8 (0.70-1.20) <br> AOR I. 14 (0.79-I.64) <br> ALL: <br> APR 0.87 (0.42-I. 8 I ) <br> AOR I.2I (0.70-2.08) <br> AOR I.I5 (0.79-I.67) <br> AML <br> AOR 2.98 (0.70-12.75) <br> AOR 0.93 (0.25-3.45) <br> AOR 0.90 (0.34-2.38) | Age 0-14 years <br> Matched for age, gender, area <br> Conditional logistic regression <br> Maternal age, mat education, household income, ethnicity, and no of residences since birth <br> Not directly adjusted maternal <br> Smoking, but maternal risk estimates did not change when paternal smoking patterns were considered |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Magnani et al. (1990) | Case control 1974-1980 198\|-1984 | 142 ALL <br> 22 AnLL <br> 19 (NHL) <br> 307 controls | Leukaemia and lymphoma <br> Paternal smoking preconception and up to childbirth: <br> ALL AOR 0.90 (0.57-1.42) <br> AnLL AOR 0.9 (0.3-2.1) <br> Lymphoma AOR 6.7 (I.0-43.4) | Personal Interview <br> Mean age: <br> Cases 6.1 (3.6) <br> Control 6.6 (3.5) <br> Four residence strata <br> Adjusted for socio-economic status | Low |
| Mattioli et al. (2014), Italy | Case control (SETIL study) 1998-2003 | 82 AnLL cases <br> 916 ALL cases <br> 1044 controls <br> (I28 matched to AnLL and 916 matched to ALL cases) | Acute non-Lymphatic Leukaemia (AnLL) <br> Paternal smoking in the conception period <br> I-I0 CPD: AOR I. 34 (0.65-2.76) <br> $\geq 11$ CPD: AOR 1.79 (I.01-3.15) | Personal interview of parents <br> Age 0-10 years Matched for date of birth, sex, residence <br> Inverse probability weighting adjusting for sex, provenience, birth order, <br> BW, breast feeding, parental educational level, age, birth year, <br> occupational exposure to benzene <br> Not directly adjusted maternal smoking but no association on AnLL and maternal smoking during pregnancy | Low |
| Menegaux et al. (2007), France | Case control 1995-1998 | 472 cases <br> 407 ALL <br> 62 AML <br> 3 other <br> 567 controls | Childhood acute leukaemia (ALL and AML) <br> Paternal smoking 3 months before pregnancy <br> All acute leukaemia <br> $\leq 20$ CPD: AOR $1.2(0.9-1.6)$ $>20 C P D:$ AOR I. $0(0.6-1.7)$ <br> ALL <br> $\leq 20$ CPD: AOR I.2 (0.9-I.6) <br> $>20$ CPD: AOR I. 2 (0.7-2.0) <br> AML <br> $\leq 20$ CPD: AOR 0.9 (0.5-I.7) <br> $>20$ CPD: AOR 0.2 (0.02-I.7) | Standardised self- administered questionnaire to mothers <br> Age < 15 years <br> Matched for age, gender, region <br> Adjusted for matched age, gender, region, socio-professional category, <br> birth order <br> Not directly adjusted for maternal smoking but not significant | Low |
| Metayer et al. (20\|3), USA | Case control (NCCLS study) 1996-2008 | 767 ALL 135 AML <br> 1139 controls | ALL and AML <br> Paternal prenatal smoking ( 3 month before and/or during pregnancy) <br> ALL: AOR I.I7 (0.91-I.50)* <br> AML: AOR I. 36 (0.82-2.24)* <br> Paternal prenatal smoking and child's passive smoking <br> ALL: AOR 0.94 (0.69-1.27)** <br> AML: AOR I.I4 (0.55-2.39)** | Phase I: Self-administered questionnaire/ <br> Phase 2: In-person interview of mainly mothers <br> Age < 15 years <br> Matched on age, maternal race, and Hispanic ethnicity <br> Adjusting for matching variables and household income <br> *Not adjusted for maternal smoking but no significant association with <br> ALL or AML <br> **adjusted for maternal prenatal smoking <br> Expansion of Chang et al. (2006) | Low |
| Milne et al. (2012), Australia | Case control (Aus-ALL study) 2003-2006 | $\begin{aligned} & 388 \text { cases } \\ & 868 \text { controls } \end{aligned}$ | ALL <br> Paternal smoking during conception year: <br> Any: AOR I. 22 (0.92-1.6I) <br> I-I4 CPD: AOR I.00 (0.66-I.52) | Self-administered questionnaires from both parents <br> Age < 15 years <br> Matched by age, sex, state of residence | Low |

## Table X Continued

| Author, year, country | Study design | Number of deliveries and children | Result <br> Outcomes (Risk estimates) | Outcomes <br> Adjustments | Quality assessment |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | > 15 CPD: AOR I. 35 (0.98-I.86) | Adjusted for matching variables, paternal age, parental education, ethnicity Maternal smoking was not associated with ALL and paternal smoking unchanged when adjusted for maternal smoking (data not shown) |  |
| Orsi et al. (2015), <br> France | $\begin{aligned} & \text { Case control (ESTELLE } \\ & \text { study) } \\ & 2010-2011 \end{aligned}$ | $\begin{aligned} & 747 \mathrm{CL} \\ & 636 \mathrm{ALL} \\ & 100 \mathrm{AML} \\ & 142 \mathrm{I} \text { controls } \end{aligned}$ | All leukaemia (AL), ALL, AML <br> Paternal preconception smoking: <br> AL: AOR I.3 (I.0-1.6) <br> ALL: AOR I. 2 (0.9-1.6) <br> AML: AOR I. 6 (I.0-2.8) <br> Paternal smoking during pregnancy: <br> AL: AOR I. 3 (I.I-I.6) <br> ALL: AOR 1.3 (I.0-1.6) <br> AML: AOR I.6 (I.0-2.5) | Telephone interview with parents, mostly mothers <br> Age <15 years <br> Matched for age, sex <br> Adjusted for age, sex, mother's age and education, birth order and maternal smoking | Low |
| Pang et al. (2003), UK | Case control (UKCCS) \|99|-96 | 3585 case fathers <br> 6987 control fathers | Leukaemia <br> Paternal preconception smoking I-I9 CPD: AOR I.I2 (0.96-I.32) 20+ CPD: AOR I.OI (0.87-I.17) <br> ALL: AOR I. 04 (0.9I-I.18) <br> AML: AOR I. 07 (0.80-I.43) | Personal interview with parents <br> Age <15 years <br> Matched for sex, age, region <br> Adjusted for matching variables, parental age, deprivation score | Medium |
| Rudant et al. (2008), France | Case control (ESCALE study) 2003-4 | 647 ALL <br> 102 AML <br> 1681 controls <br> 128 HL <br> 848 controls <br> 164 NHL <br> 1312 controls | Hematopoietic malignancies <br> Paternal smoking from the year prior to the child's birth to the interview <br> ALL: AOR I. 4 (I.I-I.7) <br> AML: AOR I.5 (I.0-2.3) <br> Hodgkin's lymphoma (HL): AOR I. 2 (0.8-I.7) <br> Non-Hodgkin’s lymphoma (NHL): AOR I. 6 (I.I-2.3) <br> <10 CPD: <br> ALL: AOR I. 2 (0.8-1.6) <br> AML: AOR I.4 (0.7-2.9) <br> HL: AOR I.4 (0.7-2.6) <br> NHL: AOR I.5 (0.8-2.6) <br> 10-I9 CPD: <br> ALL: AOR I. 2 (0.9-I.6) <br> AML: AOR I.3(0.7-2.4) <br> HL: AOR 0.8 (0.4-1.6) <br> NHL: AOR I.7 (I.I-2.7) <br> 20+ CPD: <br> ALL: AOR I. 7 (I.3-2.I)* <br> AML: AOR I.7(I.0-2.9)** <br> HL: AOR I. 2 (0.7-2.0) <br> NHL: AOR I.7 (I.I-2.6)*** | Telephone interview of mothers <br> Age < 15 years <br> Matched for age, gender <br> Adjusted for age. Gender, parental professional category, maternal age at the time of birth <br> Maternal smoking was not associated with significant increased risk <br> Trend analyses: $\begin{aligned} & * P<0.0001 \\ & * * P<0.045 \\ & * * * P<0.01 \end{aligned}$ | Low |
| Schuz et al. (I999), Germany | Case control <br> (NW and NI study) | $\begin{aligned} & 2354 \text { cases } \\ & 2588 \text { controls } \end{aligned}$ | Acute leukaemia and NHL <br> Paternal smoking before pregnancy | Questionnaire followed by telephone interview by parents Age < I 5years | Medium |


|  | NW: 1992-97 <br> NI: 1980-94 | 955 Acute leukaemia <br> 955 controls <br> 221 NHL <br> 2540 controls | Acute leukaemia (ALL and AnLL) I-I0 CPD AOR I.I (0.8-I.5) II-20 CPD AOR I.0 (0.8-।.2) >20 CPD AOR 0.9 (0.7-I.2) NHL I-I0 CPD AOR I.6 (I.0-2.5) I I-20 CPD AOR I.I (0.7-I.6) >20 CPD AOR I.I (0.7-I.8) |
| :---: | :---: | :---: | :---: |
| Shu et al. (1996), USA | Case control 1983-88 <br> (CCG study) | 302 cases <br> 203 ALL <br> 88 AML <br> II other leukaemia 558 controls Paternal smoking: 191 ALL 79 AML | ALL and AML <br> Only paternal smoking I month prior to pregnancy (A) and during pregnancy (B) <br> A: <br> ALL: AOR I. 56 (I.03-2.36) <br> I-IO CPD AOR 2.40 (I.00-5.72) <br> I I-20 CPD AOR I. 33 (0.79-2.34) <br> >20 CPD AOR I.5I (0.82-2.77) <br> AML: AOR 0.75 (0.35-1.62) <br> I-IO CPD AOR 0.42 (0.09-I.95) <br> I I-20 CPD AOR 0.73 (0.27-I.94) <br> >20 CPD AOR I. 29 (0.44-3.74) <br> B: <br> ALL: AOR I. 45 (0.95-2.19) <br> AML: AOR 0.82 (0.38-1.78) |
| Sorahan et al. (200I), UK | Case control (OSCC study) 1980-83 | ```555 cases 555 controls (GP) Cases/controls: 7/9 18/16 36/35 9/5 12/3``` | ALL <br> Paternal smoking before the pregnancy $<10$ CPD: $\text { GP: } 0.99 \text { (0.35-2.85) }$ 10-19 CPD $\text { GP:I. } 34 \text { (0.62-2.91) }$ <br> 20-29 CPD <br> GP: I. 32 (0.72-2.45) <br> 30-39 CPD <br> GP: 2.33 (0.7I-7.63) <br> 40+ CPD <br> GP: 5.29 (1.3I-2I.30) <br> $P$ for trend $P=0.06$ |
| Other cancers $\boldsymbol{n}=19$ |  |  |  |
| Barrington-Trimis et al. (2013), USA | Case control \|984-199| | 202 cases <br> 286 controls <br> Only paternal smoking: <br> 25 cases <br> 27 controls | Brain tumours <br> Only paternal smoking during pregnancy AOR I. 24 (0.66-2.35) |
| Bunin et al. (I994), USA | Case control 1986-1989 | $\begin{aligned} & \text { I55 AP } \\ & \text { I66 PNET } \\ & 32 \text { I Controls } \\ & 64 / 63 \\ & 60 / 58 \\ & 86 / 82 \\ & 85 / 88 \end{aligned}$ | Astrocytic gliomas (AG) <br> Primitive neuroectodermal tumours (PNET) <br> Paternal smoking during pregnancy <br> AG: AOR I. 0 (0.6-I.7) <br> PNET: AOR I.0 (0.6-I.7) <br> Paternal smoking ever <br> AG: AOR I.I (0.7-I8.0) <br> PNET: AOR 0.9 (0.6-I.5) |
|  | SEARCH program | 1218 cases | Brain tumours |

Matched for gender, age, region
Adjusted for socio-economic status
Not adjusted for maternal smoking, but no association with maternal smoking
Study also includes estimates on CNS tumours, neuroblastoma,
nephroblastoma, bone tumour, soft tissue sarcoma and no associations was found

Telephone interview with mothers and fathers (71\%)
Age $\leq 18$ months
Matched by age, region.
Adjusted for sex, paternal age, education, maternal alcohol consumption during pregnancy
Maternal smoking I month prior to pregnancy and during pregnancy was not associated with increased risk of ALL or AML

Interview of parents
Child age < 15 years
Matched on region, sex, date of birth
Adjusted for maternal age, paternal age, SES, ethnicity

## In-person maternal interview

Age $\leq 10$ years
Matched by age, sex, study centre
Adjusted for race, sex, age at diagnosis, maternal education, birth year,
centre
Trained interviewers with parents
Child age <6 years
Matched on race, year of birth, telephone area code and prefix
AG: Adj. income level
PNET: No adjustment

Nine centres in 7 countries

## Table X Continued

| Author, year, country | Study design | Number of deliveries and children | Result <br> Outcomes <br> (Risk estimates) | Outcomes Adjustments | Quality assessment |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Filippini et al. (2002), Italy | 1980-1992 | $\begin{aligned} & 2223 \text { controls } \\ & 633 / 1190 \end{aligned}$ | AOR I.I (0.9-1.2) | In-person interview of mostly mothers <br> Child age 0-19 years <br> Post hoc strata matched on age, sex and centre <br> Adjusted for matched variables and maternal level of education |  |
| Gold et al. (I993), USA | Case control \|977-8| | 361 cases <br> 1083 controls <br> Only paternal smoking: <br> 81 cases <br> 247 controls | Brain tumours <br> Only paternal smoking during year index child was born: <br> AOR 0.94 (0.66-1.33) | Structured interview from each parent <br> Age <18 years <br> Matched for age, sex, maternal race <br> Cases represent $85 \%$ of cases identified by the registries | Medium |
| Hu et al. (2000), China | Case control 1991-1996 | 82 cases <br> 246 controls <br> Paternal smoking + no maternal smoking: <br> 44 cases <br> 124 controls | Brain tumours <br> Smoking PY <br> AOR I.I6 (0.65-2.08) | During hospitalization, paternal and maternal interviews by trained interviewers <br> Age < 19 years <br> Matched for sex, age, area of residence <br> Adjusted for maternal education, family income | Low |
| Ji et al. (I997), <br> China | Case control \|98|-|99| | $\begin{aligned} & \text { \|98।-9\| } \\ & 642 \text { cases } \\ & 642 \text { controls } \\ & \text { Brain tumours } \\ & \text { 107 pairs } \\ & \text { Acute leukaemia, } \\ & 166 \text { pairs } \\ & \text { Lymphoma } \\ & 87 \text { pairs } \end{aligned}$ | Brain tumours <br> Paternal smoking before conception <2 PY <br> All cancers: AOR I.2 (0-8-I.8) <br> Brain tumours: AOR I.5 (0.5-4.4) 2-5 PY <br> All cancers: AOR I.3 (0.9-2.0) <br> Brain tumours: AOR I.7 (0.5-5.8) <br> >5 PY prior to conception <br> All cancers: AOR I.7 (I.2-2.5) <br> Brain tumours: AOR 2.7 (0.8-9.9) | Paternal and maternal interviews by trained interviewers <br> Age <15 years <br> Matched for sex, year of birth <br> Adjusted for BW, income, paternal age, education and alcohol | Low |
| John et al. (1991), USA | Case control 1976-83 | 1976-1983 <br> 223 cases <br> 196 controls <br> 60 exposed cancers <br> 45 exposed controls | Brain tumours <br> Paternal smoking in preconception period in the absence of maternal smoking <br> Brain tumours: AOR I.6 (0.7-3.5) <br> All cancers: AOR I.2 (0.8-2.I) | Personal interview <br> Prenatal exposure <br> Matched for age, sex, area <br> Absence of maternal smoking: <br> Adjusted for father's education | Low |
| Johnson et al. (2013)USA | Case control 2000-2008 (Cases) <br> 1994-2008 (controls) | 383 cases <br> 387 controls <br> A: Paternal smoking within the year before pregnancy: <br> II 5 cases, 84 controls <br> B: Paternal smoking during pregnancy 95 cases, 69 controls | Hepatoblastoma <br> Paternal smoking the year before pregnancy <br> A: AOR I. 4 (I.0-2.0) <br> B: AOR I.4 (0.9-2.0) | Maternal telephone interviews <br> Age $<6$ years <br> Matched for BW, gender, birth year and region <br> Adjusted for BW, year of birth, sex, maternal race and education Not directly adjusted for maternal smoking had no influence and therefore not adjusted for | Medium |
| McCredie et al. (1994), Italy + Australia | Case control Population-based 1985-1989 | 82 cases <br> 164 controls <br> Ever smoking <br> 23 cases, 28controls <br> During pregnancy <br> 4I cases, 49controls | Malignant brain tumours <br> Paternal smoking ever (at least 3 months at any time before pregnancy) <br> AOR 2.0 (I.0-4.I)* <br> Paternal smoking during pregnancy <br> AOR 2.2 (1.2-3.8) | Structured at home interviews with mothers <br> Child age up to 14 years <br> Matched on sex and age <br> *Ever smoked is adjusted for fathers schooling | Low |


| Milne et al. (2013), Australia | (Aus-CBT study) | 941 controls | Paternal smoking preconception | Age <15 years |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  | 2005-2010 | Preconception: | AOR 0.99 (0.71-1.38) | Matched for age, sex, state of residence |  |
|  |  | 74 cases | I-I4 CPD: AOR 1.31 (0.82-2.11) | Adjusted for matching variables, ethnicity, year of birth group, parental |  |
|  |  | 222 controls | I5+ CPD: AOR 0.83 (0.55-1.24) | age, household income |  |
|  |  | During pregnancy | Paternal smoking during pregnancy* | *Results shown are not adjusted for maternal smoking, but no |  |
|  |  | 71 cases | AOR I. 04 (0.74-I.46) | association was found with maternal smoking |  |
|  |  | 202 controls | $\begin{aligned} & \text { I-I4 CPD: AOR } 1.30(0.79-2.13) \\ & \text { I5+ CPD: AOR } 0.92(0.61-1.38) \end{aligned}$ | Similar results when analysis was restricted to children whose other parent did not smoke (data not shown) |  |
| Norman et al. (I996), USA | Case control | 540 cases | Childhood brain tumours | In-person or telephone interviews of mothers and fathers (77\%) | Medium |
|  | Population-based | 801 controls | Ever smoked AOR I.I (0.84-1.3) | Child age <20 years |  |
|  | 1984-1991 | Ever smoked: | Paternal smoking during pregnancy and no | Matched on birth year, sex, age at diagnosis |  |
|  |  | 262 cases, 380 controls | maternal smoking | Adj. matching criteria + maternal race/ethnicity |  |
|  |  | During pregnancy: 174 cases, 238 controls | AOR I.2 (0.95-1.6) |  |  |
| Pang et al. (2003), UK | Case control | 3585 case fathers | Cancer | Personal interview with parents | Medium |
|  | (UKCCS) | 6987 control fathers | Paternal smoking during the year before birth | Age <15 years |  |
|  | 1991-1996 | Paternal preconception | All cancers | Matched for sex, age, region |  |
|  |  | smoking | I-19 CPD: AOR I.II (0.98-I.25) | Adjusted for matching variables, parental age, deprivation score |  |
|  |  | All cancers | 20+ CPD: AOR I.01 (0.90-1.12) |  |  |
|  |  | 583 cases, 1003 controls | CNS tumours |  |  |
|  |  | 757 cases, 1440 controls | I-19 CPD: AOR I. 08 (0.85-1.38) |  |  |
|  |  | CNS tumours | 20+ CPD: AOR 1.03 (0.82-1.28) |  |  |
|  |  | 101 cases, 1003 controls |  |  |  |
|  |  | 138 cases, 1440 controls |  |  |  |
| Plichart et al. (2008), France | Case control | 209 cases | CNS tumours | Maternal telephone interview | Low |
|  | (ESCALE study) | 1681 controls | Only paternal smoking in the year prior the child's birth <br> AOR I.3 (I.0-I.9) | Age <15 years |  |
|  | 2003-2004 | Paternal smoking + nonsmoking mother |  | Matched for age, sex and number of children <15 years of age in the household |  |
|  |  | 74 cases, 516 controls |  | Adjusted for age, gender |  |
|  |  |  |  | No association between maternal smoking during pregnancy and CNS tumours. |  |
| Sorahan et al.(I997a), UK | Case control | 1549 cases | Death of childhood cancer | Interview parents, usually mothers (response rate 88\%) | Medium |
|  | (OSCC study) | 1549 controls | Paternal smoking at death of child, father only | Matched for sex, date of birth and region |  |
|  | 1953-1955 | 655 cases, 618 controls | ARR I. 30 (1.10-1.53) | Adjusted for social class, parental age at birth, sib-ship position, obstetric radiography |  |
| Sorahan et al. (I997b), UK | Case control | 2587 cases | Death of childhood cancer | Interview of parents, usually mothers | Medium |
|  | (OSCC study) | 2587 controls | Paternal smoking at death of child, father only | Child age $<16$ years |  |
|  | 1971-1976 | 630 cases | 14\% of the cancers could be related to | Matched for sex, date of birth, region |  |
|  |  | 573 controls | paternal smoking | Adjusted for social class, parental age at birth, sib-ship position, |  |
|  |  |  | (all cancer and onset at all ages) | obstetric radiography |  |
|  |  |  | ARR 1.29 (1.10-1.51) |  |  |
| Sorahan et al. (200I), UK | Case control | 555 cases | Childhood cancer | Interview of parents | Low |
|  | (OSCC study) | 555 controls (hospital) | Paternal smoking before the pregnancy | Child age < 15 years |  |
|  | 1980-83 | 555 controls (GP) | ARR: | Matched on region, sex, date of birth |  |
|  |  | Cases/GP/Hospital | <10 CPD: | Adjusted for maternal age, paternal age, SES, ethnicity |  |
|  |  | 26/34/27 | GP: 0.94 (0.53-I.66); Hospital: 0.92 |  |  |
|  |  | 79/60/70 | (0.5I-I.65) |  |  |
|  |  | 114/122/121 | 10-19 CPD |  |  |
|  |  | 23/32/48 |  |  |  |

## Table X Continued

| Author, year, country | Study design | Number of deliveries and children | Result <br> Outcomes (Risk estimates) | Outcomes <br> Adjustments | Quality assessment |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | 28/21/40 | GP: I. 63 (I.10-2.4I); Hospital: I. 06 <br> (0.72-1.56) <br> 20-29 CPD <br> GP: I. 46 (I.05-2.03); Hospital: I. I I <br> (0.80-1.53) <br> 30-39 CPD <br> GP: 0.95 (0.52-I.73); Hospital: 0.45 <br> (0.26-0.77) <br> 40+ CPD <br> GP: I. 77 (0.94-3.34); Hospital: 0.66 <br> (0.39-I.II) <br> $P$ for trend <br> GP $P=0.02$; Hospital $P=0.16$ <br> CNS tumours also stratified on CPD, but no total ARR <br> $P$ for trend 0.67 <br> Data adjusted for maternal smoking is not shown but <br> with a significant positive trend $(P=0.03)$ <br> between cancer <br> and paternal smoking compared to GP controls |  |  |
| Sorahan and Lancashire (2004), UK | Case control (OSCC study) <br> Deaths 1953-55 <br> 1971-76 <br> \|977-8| | 5777 case control <br> matched pairs <br> Paternal smoking only: <br> All cancers <br> 1637 cases <br> 1545 controls <br> Hepatoblastoma <br> 43 cases <br> 8 cases <br> \| 545 controls | Death of childhood cancer and hepatoblastoma <br> All cancers <br> ARR 1.28 (I.15-I.42) <br> Hepatoblastoma <br> ARR I. 23 (0.46-3.28) | Interview parents, usually mother <br> Child age < 16 years <br> Matched for sex, age at death, year of death <br> Adjusted for sex, age at death, year of death, social class, sib-ship position, maternal age, paternal age, obstetric radiography | Medium (all cancers) <br> Low (hepatoblastoma) |
| Schuz et al. (1999), Germany | Case control (NW and NI study) <br> NW: 1992-97 <br> NI:1980-94 | NW: 1992-97 <br> NI:1980-94 <br> 2358 cases <br> 2588 controls <br> 385 CNS tumours <br> I55 neuroblastomas <br> 2540 nephroblastomas <br> 95 bone tumours <br> 133 soft tissue sarcomas | CNS tumour, Neuroblastoma, <br> Nephroblastoma, Bone tumour, Soft tissue sarcoma <br> Paternal smoking before pregnancy <br> I-IO CPD: <br> CNS tumour: AOR 0.8 (0.5-I.2) <br> Neuroblastoma: AOR 0.6 (0.3-1.1) <br> Nephroblastoma: AOR 0.8 (0.4-I.4) <br> Bone tumour: AOR 0.5 (0.2-1.2) <br> Soft tissue sarcoma: AOR 0.8 (0.4-1.6) <br> II-20 CPD: <br> CNS tumour: AOR I.I (0.8-I.4) <br> Neuroblastoma: AOR I.I (0.7-1.6) | Questionnaire followed by telephone interview by parents <br> Age < 15 years <br> Matched for gender, age, region <br> Adjusted for socio-economic status <br> Not adjusted for maternal smoking, but no association with maternal smoking. <br> Study also includes estimates on <br> Acute leukaemia and NHL | Low |

Nephroblastoma: AOR 0.8 (0.5-1.3)
Bone tumour: AOR 0.8 (0.4-1.3)
Soft tissue sarcoma: AOR I.2 (0.8-I.8)
$>20 \mathrm{CPD}$
CNS tumour: AOR I.0 (0.7-I.4)
Neuroblastoma: AOR I. 2 (0.7-2.1)
Nephroblastoma: AOR 0.9 (0.5-I.6)
Bone tumour: AOR 0.9 (0.4-1.8)
Soft tissue sarcoma: AOR 0.9 (0.4-I.6

| Yang et al. (2000), | Case control |
| :--- | :--- |
| USA \& Canada | (CCG and POG |
|  | studies) | studies)

1992-94

504 cases 504 controls
Preconception
137 cases, 122 controls

Neuroblastoma
Paternal smoking one month before conception
AOR I.2 (0.8-I.6)

## Cardio-metabolic outcomes $(\mathbf{n}=9)$

| Brion et al. (2007), UK | Cohort study Avon longitudinal study | 6396 children (Model I) <br> 3736 children (Model 5) | Blood pressure at 7 years <br> Systolic blood pressure: <br> Model I: Beta $0.44(-0.07-0.95) P=0.09$ <br> Model 5: Beta $0.17(-0.52-0.86) P=0.6$ <br> Diastolic blood pressure: <br> Model I: Beta $0.10(-0.26-0.47) P=0.6$ <br> Model 5: Beta $-0.25(-0.72-0.22) P=0.3$ | Questionnaires sent to partners at 18 weeks gestation on if they had smoked regularly in the last 9 months <br> Model I: Child age, sex <br> Model 5: Additionally, adjusted for maternal/partner factors, social factors, breast feeding | Medium |
| :---: | :---: | :---: | :---: | :---: | :---: |
| de Jonge et al. (20\|3), US | Nurse's Health Study II and Nurses' Mothers' cohort 1989-2007 | 5777 non-smoking mothers 3078 paternal smoking 2699 no paternal smoking | Hypertension in daughters in adulthood (self-reported physician diagnosed) Paternal smoking during pregnancy Maternal age: ARR I.12 (1.06-1.18) + perinatal variables: ARR 1.09 (I.03-I.15) + BW: ARR I. 08 (I.03-I.14) + adult life variables: ARR 1.08 (I.02-1.14) + body shape and weight until age: ARR 18:I. 07 (1.01-1.13) + current BMI: I. 04 (0.99-I.I0) | Self-administered questionnaires to nurse's mothers 2001 Cox proportion hazard models Multiple adj. and additional adj. for perinatal variables, adult life variables, body shape and weight until age 18 years, current BMI | Medium |
| Durmus et al. (201I), The Netherlands | Prospective cohort study 2002-2006 | 4028 non-Smoking <br> mothers <br> Paternal smoking during <br> pregnancy: <br> 1397 fathers <br> 0-4 CPD: 607 fathers <br> $\geq 5$ CPD: 753 fathers <br> 2572 no paternal smoking | BMI at $3,6,12,24,36,48$ months <br> Paternal smoking during pregnancy and difference in BMI at 12 months: <br> Standardized coefficients ( $95 \% \mathrm{Cl}$ ): $0.06(-0.0 \mathrm{I}, 0.13)$ <br> $0-4$ CPD $0.04(-0.05,0.13)$ <br> $\geq 5$ CPD $0.08(-0.01,0.17)$ <br> $P$ for trend $P=0.01$ <br> Similar no difference in BMI at $3,6,24,36$ and 48 months and no trend | Postal questionnaires to mothers <br> Linear mixed models <br> Adj. Child's age at visit, sex, paternal ethnicity and education, paternal height and weight and breast feeding (yes/no) <br> Reporting bias <br> Similar information completed by the fathers in 3358 participants - good agreement between mat and pat assessment | Medium |
| Florath et al. (2014), Germany | Prospective cohort Born 2000-2001 | 609 healthy mature newborns | BMI at child age 8 years Smoking during pregnancy BMI at 8 years | During hospitalization after delivery standardized maternal interviews by trained interviewers. Follow-up to age 8 years Linear regression | Medium |

Table X Continued

| Author, year, country | Study design | Number of deliveries and children | Result <br> Outcomes <br> (Risk estimates) | Outcomes <br> Adjustments | Quality assessment |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Paternal smoking during pregnancy in nonsmoking mothers I57 paternal smoking 401 no paternal smoking | Adjusted regression coefficient: 0.34 (0.0I-0.66) | Adjusted for paternal BMI and education, maternal pre-pregnancy BMI, BW, monthly weight gain, exclusive breast feeding, body height, TV consumption, sports activities, diet score at 8 years and age at anthropometric measurements. <br> Conclusion: <br> Residual confounding conditions in smoking families by living rather than specific intrauterine exposure may account for the increased risk of offspring overweight |  |
| Howe et al. (2012), UK | Cohort study <br> Avon longitudinal study | Height: <br> 4832 children <br> PI: 4777 children <br> BMI: 4534 children | Growth 29-120 months <br> Girls: 0.0012 ( 0.002 I ), $P=0.0 \mathrm{I}$ <br> Boys: -0.00 I 2 ( 0.0020 ), $P=0.8$ <br> Ponderal index 2-24 months: <br> Girls: 0.0043 ( 0.0078 ), $P=0.35$ <br> Boys: -0.00 I I ( 0.0069 ), $P=0.73$ <br> BMI I03-I20 months: <br> Girls: 0.0042 ( 0.0036 ), $P=0.54$ <br> Boys: 0.0033 ( 0.002 I), $P=0.77$ | Self-reported data <br> Height 0-10 years <br> Ponderal index 0-2 years <br> BMI 2-10 years <br> Maternal education, household social class, parity, maternal age, maternal height, maternal BMI, gestational age, breast feeding | Medium |
| Kwok et al. (20I0), Hong Kong | Birth Cohort study 1997 | Non-smoking mothers: <br> 7924 children <br> 6710 children with BMI at <br> 7 years <br> 6519 children with BMI at <br> II years of age | BMI and height at child age 7 and 11 years Daily prenatal and early postnatal paternal smoking: <br> BMI, Z-score difference, mean ( $95 \% \mathrm{Cl}$ ) <br> Child age 7 years: 0.10 (0.02-0.19) <br> Child age II years: 0.16 (0.07-0.26) <br> No difference in height $Z$ scores | Standardized self- administered questionnaire at maternal and childhealth centres <br> Daily prenatal and early postnatal paternal smoking in non-smoking women <br> Adjusted for gender, birth order, highest parental education, mother's place of birth, pubertal status (for II years) highest parental occupation, household income per person, breast feeding history, number of hospital admissions attributable to infections at 0 to 6 months | Medium |
| Leary et al. (2006), UK | Cohort study Avon longitudinal study | Examination at 9 years 6470 children 5615 children* <br> 3649 children** | BMI, total fat, truncal fat, total lean (DXA scanner) at mean child age 9.9 years Paternal smoking during pregnancy <br> *BMI: beta 0.1 I ( $0.05,0.17$ ) $<0.00$ I <br> *Total fat: beta $0.08(0.03-0.13) P=0.00$ I | Questionnaires to mothers <br> Adjusted for maternal smoking <br> *Sex, child age at DXA-scan, ${ }^{* *}$ Additionally adjusted for maternal, partner, social and infant feeding factors | Medium |
| Taal et al. (2013), The Netherlands | Prospective cohort study 2008-2012 | Non-smoking mothers during singleton pregnancy 4070 cases <br> Fathers smoking 1298 <br> cases <br> Fathers non-smoking 2369 controls | Stroke, volume, cardiac output, larger AOD (aortic root diameter), fractional shortening at child age 6 years <br> Regression coefficients <br> Mean Systolic blood pressure ( mmHg ) <br> -0.18 (-0.69-0.33) test for trend over <br> smoking cat. 0.74 I <br> Mean diastolic blood pressure ( mmHg ) <br> $0.12(-0.39-0.52)$ test for trend over smoking <br> cat. 0.702 <br> Aortic root diameter (mm) <br> Difference 0.17 (0.05-0.28) | Questionnaires in second and third trimester Mixed models and multiple linear regression models Adjusted for maternal age, parity, mixed educational level, prepregnancy $\mathrm{BMI}, \mathrm{BP}$ at intake, sex, GA, BW, breast feeding status, current age and BMI | Medium |



| Author, year, country | Study design | Number of deliveries and children | Result <br> Outcomes <br> (Risk estimates) | Outcomes Adjustments | Quality assessment |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Zhu et al. (2014), <br> Denmark | Birth cohort and discharge diagnosis from Medication Registry | 50870 mothers participated in 7-years questionnaire 14004 singletons with paternal smoking only 360 (2.6\%) singletons with ADHD <br> Both non-smokers ADHD: $\text { 892/49 } 072 \text { (।.8\%) }$ | ADHD <br> Paternal smoking with no maternal smoking AHR I. 29 (I.14-I.47) | Questionnaire during pregnancy and at follow-up 7- years of age (mother) <br> Cox regression adjusted for maternal age, parity, alcohol, SES, psychopathology, sex, diagnosis, education (registry) | Medium |

The systematic literature search revealed a huge number of articles which were scrutinized, and 238 of these publications were selected for inclusion. Although the quality of included articles varied, several large cohort studies of high quality were identified.

A majority of the included publications investigated the effect of paternal age on the health of children. A previous systematic review concerning the effect of paternal factors on obstetric outcomes found that extremes in paternal age were associated with an increase in LBW, and there was also an association between paternal height and BW of offspring (Shah, 2010). Another recent report summarized the association between paternal age and health of offspring in a narrative way (Nybo Andersen and Urhoj, 2017). They reported a strong association between paternal age and some specific congenital syndromes, namely cleft palate, acute lymphatic leukaemia, ASD and schizophrenia.

While no clear definition seems to exist for advanced paternal age, many studies have used 40 years and above as an age limit. A common problem for many studies of paternal factors, particularly paternal age, is the strong and well-known confounding factor, namely the effect of maternal age on obstetric and child outcome. A similar influence is true for other exposures, such as smoking and BMI. In our meta-analyses we therefore only included publications which had adjusted for maternal age or smoking, or where it was clear that the reference group was young mothers or non-smoking mothers, respectively. We also included some studies in which univariate analysis of maternal smoking was insignificant, and thus did not adjust for maternal smoking in the multivariate analyses.

## Paternal age

Among obstetric outcomes, we found a small but significantly increased risk of stillbirth associated with paternal age. We also found significantly higher risks of birth defects, and specifically of orofacial clefts and trisomy 21 . For other obstetric outcomes and selected birth defects we could not identify any increased risks. For gastroschisis, there seems to be evidence of an increased risk associated with younger fathers (Kazaura et al., 2004a, Archer et al., 2007, Yang et al., 2007, Materna-Kiryluk et al., 2009). The mechanism behind such an association is suggested to be of socio-economic origin, with certain life-style factors more common among young fathers (drugs, smoking, etc) while the increase in risk associated with higher paternal age is possibly of genetic origin, depending on a higher frequency of de novo mutation in sperm of older fathers.

Children born to older fathers have been reported as having a higher risk of various cancer types (Sartorius and Nieschlag, 2010). A meta-analysis (Sergentanis et al., 2015) reported an increased risk of childhood leukaemia associated with higher parental ages. For the risk of ALL, they reported an association with both increased maternal and paternal age. In our meta-analysis we did not find an association between ALL and advanced paternal age. A possible explanation for our result could be that we only included studies that adjusted for maternal age. For other cancer types, it was not possible to carry out meta-analysis, either because the studies were too few, of too low quality, or did not adjust for maternal age.

Psychiatric disorders and diseases like autism/ASD (Hultman et al., 2011; Wu et al., 2017) and schizophrenia (Miller et al., 2010) have been associated with advanced paternal age. From earlier studies, the risk seems to increase linearly without any particular threshold. This


Figure 19 Forest plot describing the association between paternal smoking and risk for brain tumours in the offspring.
link with advanced paternal age has been found in studies from different countries although the magnitude of the associations varies.

ASD is a chronic disease and includes different conditions such as infantile autism, Asperger's syndrome, atypical autism and pervasive development disorder. There are different theories explaining the aetiology. The genetics of ASD seem complex and may involve genetic, epigenetic and environmental factors (Waye and Cheng, 20I7). The mean number of de novo mutations in human spermatozoa has been found to increase by around two per year (Kong et al., 2012). This increased frequency of de novo mutations of older men may result in both gain and loss of DNA region copy numbers, and may explain some of the effects associated with increased paternal age. Other explanations for the paternal effects may include the fact that these men are more often in the higher or lower socio-economic groups, they are more likely to be overweight and obese, and they are more likely to smoke and have a higher alcohol intake. Furthermore, these men more often suffer from diverse physical and mental health problems (Nilsen et al., 2013).

In our meta-analysis of autism, 16 studies were included and a higher risk of autism/ASD was associated with increasing paternal
age (pooled estimate $1.25,95 \% \mathrm{Cl}$ I.20-I.30). All studies adjusted for maternal age. This finding is in line with previous meta-analyses (Hultman et al., 201I; Wu et al., 2017).

Our meta-analysis of schizophrenia included II original articles, all of these studies adjusted for maternal age. A higher risk of schizophrenia was observed in the children of older men (pooled estimate I.3I, 95\% Cl I.24-I.38). The previous meta-analysis by Miller et al. (2010) found a higher level of risk in the oldest fathers ( $\geq 50$ years), of AOR 1.66 (95\% CI I.46-I.89). Similarly, the meta-analyses by Torrey et al. (2009) and Wohl and Gorwood (2007) found a strong association between advanced paternal age and an increased risk of schizophrenia in offspring. Limitations in some of the studies include few outcomes of interest, and poorly defined classification of diagnosis. Another limitation is that it is not possible to adjust for genetic and environmental factors that could be important confounders in the association between advanced paternal age and adverse outcomes in offspring. Furthermore, there is a heterogeneity of different age categories that complicates comparison between studies. Our results are in line with previous studies and indicate an association between advanced paternal age and increased risk of autism, schizophrenia and other psychiatric diseases.

Table XI Summary results of the meta-analyses of the association between paternal factors and perinatal and paediatric outcomes.

| Exposure | Outcome | Pooled estimate (with 95\% CI) | Certainty of evidence GRADE |
| :---: | :---: | :---: | :---: |
| Paternal age | PTB | 1.02 (1.00-1.05) | $\oplus \oplus \bigcirc \bigcirc$ |
|  | Low BW | 1.00 (0.97-1.03) | $\oplus \oplus \bigcirc \bigcirc$ |
|  | Stillbirth | 1.19 (1.10-1.30) | $\oplus \oplus \bigcirc \bigcirc$ |
|  | Children with any birth defects | 1.05 (1.02-1.07) | $\oplus \oplus \oplus \bigcirc$ |
|  | CHDs | 1.03 (0.99-1.06) | $\oplus \oplus \oplus \bigcirc$ |
|  | Orofacial clefts | 0.99 (0.95-1.04) | $\oplus \oplus \bigcirc \bigcirc$ |
|  |  | 1.14 (1.02-1.29)* |  |
|  | Gastroschisis | 0.88 (0.78-1.00) | $\oplus \oplus \oplus \bigcirc$ |
|  | Spina bifida | 0.97 (0.90-1.04) | $\oplus \oplus \oplus \bigcirc$ |
|  | Trisomy 21 | 1.13 (1.05-1.23) | $\oplus \oplus \oplus \bigcirc$ |
|  | Acute | 1.08 (0.96-1.2I) | $\oplus \oplus \oplus \bigcirc$ |
|  | lymphoblastic |  |  |
|  | leukaemia |  |  |
|  | Autism and ASDs | 1.25 (1.20-1.30) | $\oplus \oplus \oplus \bigcirc$ |
|  | Schizophrenia | 1.31 (1.23-1.38) | $\oplus \oplus \oplus \bigcirc$ |
| Paternal BMI | No meta-analysis |  |  |
| Paternal | PTB | 1.16 (1.00-1.35) | $\oplus \oplus \bigcirc \bigcirc$ |
| smoking | Low BW | 1.10 (1.00-1.21) | $\oplus \oplus \bigcirc \bigcirc$ |
|  | SGA | 1.22 (1.03-1.44) | $\oplus \oplus \bigcirc \bigcirc$ |
|  | CHDs | 1.75 (1.25-2.44) | $\oplus \oplus \bigcirc \bigcirc$ |
|  | Orofacial clefts | 1.51 (1.16-1.97) | $\oplus \oplus \bigcirc \bigcirc$ |
|  | Brain tumours | 1.12 (1.03-1.22) | $\oplus \oplus \bigcirc \bigcirc$ |

*Exposure: Paternal age $>45$ years.

Paternal, BMI, height and/or weight. There are only limited data on the impact of paternal obesity at the time of conception on short and long-term health outcomes for children. Information on the father's weight and BMI was often documented after the birth of the child, and those studies were excluded from the analysis. Paternal obesity has been connected to infertility, a reduced rate of live birth per cycle in ART, and increased risk of pregnancy non-viability (Campbell et al., 2015). Obese men have an increased amount of sperm with low mitochondrial membrane potential, DNA fragmentation, and abnormal morphology, all of which may have harmful effects on fertility (Campbell et al., 2015). However, if the pregnancy starts, prepregnancy paternal BMI does not seem to exert any independent effect on the risk of PTB or SGA (Mutsaerts et al., 2014). Actually, with regard to short-term outcomes, only the height of the father correlated significantly with the BW of the offspring. On the other hand, paternal anthropometrics at the time of the child's birth were associated with childhood BMI, weight and/or fat mass. However, paternal height and weight were usually either self-reported, or reported by the mother, and not measured, which increases the risk of bias. The majority of studies reported a stronger effect of maternal BMI than paternal BMI. In one study, Patro and co-workers systematically evaluated the associations of offspring BMI, or adiposity, with pre-pregnancy BMI (or adiposity) of the mother and the father (Patro et al., 2013). They hypothesized that the intrauterine environment is an independent factor in obesity development, and thus the maternal
effect is likely to be stronger ('foetal overnutrition hypothesis'), but found only limited evidence to support the hypothesis.

Paternal smoking. Our meta-analyses on short-term outcomes demonstrated a small but significant increased risk of SGA if fathers smoked, but non-significant increased risks of PTB and LBW. While the effect of maternal smoking on obstetric outcomes is evident, the effect of paternal smoking on obstetric outcomes is still equivocal. Paternal smoking significantly increased CHD and orofacial clefts, with pooled estimates of 1.75 and 1.50 respectively. The occurrence of CHD and orofacial clefts is a result of the interaction of genetic and environmental risk factors and cigarette smoke comprises numerous chemical carcinogens. These chemicals have a teratogenic effect on oocytes and sperm DNA or interfere with foetal cardiac development and other foetal structures (Deng et al., 2013; Figueiredo et al., 2015). The existing literature is not able to clarify if the increased risk of CHD associated with paternal smoking is due to a direct effect on sperm, or the result of passive smoking on oocytes or the foetus: an effect of the latter should be weaker than that of maternal smoking. The effect of maternal smoking on orofacial cleft risk is relatively well demonstrated, but an effect on cardiovascular defects is more dubious or perhaps restricted to only some forms. Thus, the effect of paternal smoking on orofacial clefts may be due to maternal passive smoking, and why the effect on CHD is more likely to be due to paternal smoking per se. Our meta-analysis of data on brain tumours showed a small but significant increased risk of brain tumours if fathers smoked, with pooled estimate of I.12. The previous meta-analyses on data on leukaemia demonstrated an association between paternal smoking at pregnancy for AML (AOR I.19-1.28) and for ALL (I.15-I.25) (Liu et al., 20। I; Milne et al., 2012; Metayer et al., 2016). Tobacco smoke is known to be leukomogenic, to introduce oxidative damage in sperm cells resulting in DNA fragmentation, and it may also cause persistent changes in miRNA (Liu et al., 20II; Milne et al., 20I2).

The association between paternal smoking and childhood cancer could hardly be due to maternal passive smoking as maternal smoking is not with certainty associated with childhood cancer. However, if the infant or young child has been exposed for passive smoking that may explain the increased cancer risk, if passive smoking has a carcinogenic effect. Most of the studies on the effects of exposure to paternal smoking during pregnancy had matched control groups, and risk estimates were adjusted for relevant confounding factors. Nevertheless, none of the studies were assessed as being of high quality. Most studies were retrospective case control studies with a limited number of cases and controls, where fathers but not mothers smoked.
In the majority of studies, the information on paternal smoking was gained from interviews with, or questionnaires from, the mothers, thus introducing recall and selection bias. Obviously, recall bias is of most concern in long-term outcomes where information on paternal smoking is obtained several years after childbirth. The extent of under reporting of paternal smoking is unknown, which may introduce some non-differential misclassification bias. Furthermore, the dissemination of information on the adverse consequences of smoking in pregnancy may have discouraged some parents from disclosing it. Even in some studies of short-term outcomes, information on BW and gestational age came in the form of maternal self-reported data rather than hospital files or national registries, meaning that accuracy is less certain. Most studies of long-term outcomes involved univariate analyses of
maternal smoking, and if not significant, the final results were not adjusted for maternal smoking. Further differentiation between the impact on long-term outcomes of paternal smoking in the preconception stage, and during and after pregnancy, is a challenge. Exposure of pregnant women to tobacco smoke in the environment may be a confounding factor and likewise, second-hand smoke after delivery is a confounder when examining long-term outcomes.

## Strengths and limitations

The major strength of this systematic review is the comprehensive literature search, identifying a huge number of relevant publications, from the beginning of the 1950s up to 2017. Another strength is the fact that it is possible to perform meta-analyses, making interpretation of the summarized literature much easier for the reader. The main limitation is the heterogeneity within the meta-analyses e.g. differences in paternal age groups, reference groups, outcome measures and statistical methods used in the studies included. This heterogeneity may, however, be of less importance since this systematic review/meta-analysis is based solely on observational studies and not on interventional trials. All estimates are thus limited to associations. Another limitation which has to be observed is that most of the estimates and $95 \% \mathrm{Cl}$ for the estimates are close to I.00. Despite adjusting for confounders, for example maternal age, residual confounders might well exist, which could explain the association between paternal age and several of the outcomes.

## Conclusion

This systematic review and meta-analysis investigating paternal age, smoking and $\mathrm{BMI} /$ weight/height as risk factors for adverse outcome in offspring, found elevated risks for selected birth defects and psychiatric disorders as well as for selected cancers and metabolic disturbances. Although these risks represent serious health effects for the children, the magnitude of these effects seems modest.

## Supplementary data

Supplementary data are available at Human Reproduction Update online.

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## Authors' roles

A.L., A.P., C.B., H.L., L.B.R., N.B.O., U.B.W. and V.S.A. contributed to the design of the study, screened articles, selected articles, performed data extraction, interpreted the data and wrote the manuscript. MP performed the statistical analyses. All authors approved the final version for submission.

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## Conflict of interest

None of the authors has any conflicts of interest to declare.

## References

Abel EL, Kruger M, Burd L. Effects of maternal and paternal age on Caucasian and Native American preterm births and birth weights. AmJ Perinatol 2002; 19:49-54.
Alio AP, Salihu HM, McIntosh C, August EM, Weldeselasse H, Sanchez E, Mbah AK. The effect of paternal age on fetal birth outcomes. Am J Mens Health 2012; 6:427-435.
Andriani H, Kuo HW. Adverse effects of parental smoking during pregnancy in urban and rural areas. BMC Pregnancy Childbirth 2014;14:414.
Archer NP, Langlois PH, Suarez L, Brender J, Shanmugam R. Association of paternal age with prevalence of selected birth defects. Birth Defects Res A Clin Mol Teratol 2007;79:27-34.
Astolfi P, De Pasquale A, Zonta LA. Late paternity and stillbirth risk. Hum Reprod 2004;19:2497-250I.
Astolfi P, De Pasquale A, Zonta LA. Paternal age and preterm birth in Italy, 1990 to 1998. Epidemiol 2006;17:218-221.
Barrington-Trimis JL, Searles Nielsen S, Preston-Martin S, Gauderman WJ, Holly EA, Farin FM, Mueller BA, McKean-Cowdin R. Parental smoking and risk of childhood brain tumors by functional polymorphisms in polycyclic aromatic hydrocarbon metabolism genes. PLoS One 2013;8:e791I0.
Basso O, Wilcox AJ. Paternal age and delivery before 32 weeks. Epidemiol 2006; 17:475-478.
Ben Itzchak E, Lahat E, Zachor DA. Advanced parental ages and low birth weight in autism spectrum disorders-rates and effect on functioning. Res Dev Disabil 2011;32:1776-1781.
Berg E, Lie RT, Sivertsen A, Haaland OA. Parental age and the risk of isolated cleft lip: a registry-based study. Ann Epidemiol 2015;25:942-947.e94I.
Bilder D, Pinborough-Zimmerman J, Miller J, McMahon W. Prenatal, perinatal, and neonatal factors associated with autism spectrum disorders. Pediatrics 2009; I23: 1293-1300.
Bille C, Skytthe A, Vach W, Knudsen LB, Andersen AM, Murray JC, Christensen K. Parent's age and the risk of oral clefts. Epidemiol 2005;16:31।-316.
Brion MJ, Leary SD, Smith GD, Ness AR. Similar associations of parental prenatal smoking suggest child blood pressure is not influenced by intrauterine effects. Hypertension 2007;49:1422-1428.
Brion MJ, Victora C, Matijasevich A, Horta B, Anselmi L, Steer C, Menezes AM, Lawlor DA, Davey Smith G. Maternal smoking and child psychological problems: disentangling causal and noncausal effects. Pediatrics 2010; I26:e57-e65.
Brondum J, Shu XO, Steinbuch M, Severson RK, Potter JD, Robison LL. Parental cigarette smoking and the risk of acute leukemia in children. Cancer 1999;85: 1380-1388.
Brown A, Bao Y, McKeague I, Shen L, Schaefer C. Parental age and risk of bipolar disorder in offspring. Psychiatry Res 2013;208:225-23।.
Brown AS, Schaefer CA, Wyatt RJ, Begg MD, Goetz R, Bresnahan MA, HarkavyFriedman J, Gorman JM, Malaspina D, Susser ES. Paternal age and risk of schizophrenia in adult offspring. Am J Psychiatry 2002;159:1528-1533.
Buizer-Voskamp JE, Laan W, Staal WG, Hennekam EA, Aukes MF, Termorshuizen F, Kahn RS, Boks MP, Ophoff RA. Paternal age and psychiatric disorders: findings from a Dutch population registry. Schizophr Res 2011;129:I28-132.
Bunin GR, Buckley JD, Boesel CP, Rorke LB, Meadows AT. Risk factors for astrocytic glioma and primitive neuroectodermal tumor of the brain in young children:
a report from the Children's Cancer Group. Cancer Epidemiol, Biomarkers Prev 1994;3:197-204.
Bunin GR, Needle M, Riccardi VM. Paternal age and sporadic neurofibromatosis I: a case-control study and consideration of the methodologic issues. Genet Epidemiol 1997; 14:507-516.
Burd L, Severud R, Kerbeshian J, Klug MG. Prenatal and perinatal risk factors for autism. J Perinat Med 1999a;27:44I-450.
Burd L, Severud R, Klug MG, Kerbeshian J. Prenatal and perinatal risk factors for Tourette disorder. J Perinat Med 1999b;27:295-302.
Byars SG, Boomsma JJ. Opposite differential risks for autism and schizophrenia based on maternal age, paternal age, and parental age differences. Evol Med Public Health 2016;2016:286-298.
Byrne M, Agerbo E, Ewald H, Eaton WW, Mortensen PB. Parental age and risk of schizophrenia: a case-control study. Arch Gen Psychiatry 2003;60:673-678.
Campbell JM, Lane M, Owens JA, Bakos HW. Paternal obesity negatively affects male fertility and assisted reproduction outcomes: a systematic review and meta-analysis. Reprod Biomed Online 2015;31:593-604.
Cardwell CR, Carson DJ, Patterson CC. Parental age at delivery, birth order, birth weight and gestational age are associated with the risk of childhood Type I diabetes: a UK regional retrospective cohort study. Diabet Med 2005;22:200-206.
Castro-Jimenez MA, Orozco-Vargas LC. Parental exposure to carcinogens and risk for childhood acute lymphoblastic leukemia, Colombia, 2000-2005. Prev Chronic Dis 2011;8:Al06.
Catalano PM, Farrell K, Thomas A, Huston-Presley L, Mencin P, de Mouzon SH, Amini SB. Perinatal risk factors for childhood obesity and metabolic dysregulation. Am J Clin Nutr 2009;90:1303-1313.
Cawley RH, Mc KT, Record RG. Parental stature and birth weight. Am J Hum Genet 1954;6:448-456.
Chang JS, Selvin S, Metayer C, Crouse V, Golembesky A, Buffler PA. Parental smoking and the risk of childhood leukemia. Am J Epidemiol 2006;163:1091-I I00.
Chen XK, Wen SW, Krewski D, Fleming N, Yang Q, Walker MC. Paternal age and adverse birth outcomes: teenager or 40+, who is at risk? Hum Reprod 2008;23: 1290-1296.
Chen YP, Xiao XM, Li J, Reichetzeder C, Wang ZN, Hocher B. Paternal body mass index (BMI) is associated with offspring intrauterine growth in a gender dependent manner. PLoS One 2012;7:e36329.
Chudal R, Gissler M, Sucksdorff D, Lehti V, Suominen A, Hinkka-Yli-Salomaki S, Brown AS, Sourander A. Parental age and the risk of bipolar disorders. Bipolar Disord 2014;16:624-632.
Chudal R, Joelsson P, Gyllenberg D, Lehti V, Leivonen S, Hinkka-Yli-Salomaki S, Gissler M, Sourander A. Parental age and the risk of attention-deficit/hyperactivity disorder: a nationwide, population-based cohort study. J Am Acad Child Adolesc Psychiatry 2015;54:487-494.e48।.
Cresci M, Foffa I, Ait-Ali L, Pulignani S, Gianicolo EA, Botto N, Picano E, Andreassi MG. Maternal and paternal environmental risk factors, metabolizing GSTMI and GSTTI polymorphisms, and congenital heart disease. Am J Cardiol 2011;108: 1625-1631.
Croen LA, Najjar DV, Fireman B, Grether JK. Maternal and paternal age and risk of autism spectrum disorders. Arch Pediatr Adolesc Med 2007;16 1:334-340.
Cross PK, Hook EB. An analysis of paternal age and 47,+21 in 35,000 new prenatal cytogenetic diagnosis data from the New York State Chromosome Registry: no significant effect. Hum Genet 1987;77:307-3I6.
Crump C, Sundquist K, Sieh W, Winkleby MA, Sundquist J. Perinatal and family risk factors for non-Hodgkin lymphoma in early life: a Swedish national cohort study. J Natl Cancer Inst 2012;104:923-930.
Crump C, Sundquist J, Sieh W, Winkleby MA, Sundquist K. Perinatal and familial risk factors for brain tumors in childhood through young adulthood. Cancer Res 2015;75:576-583.
D'Onofrio BM, Rickert ME, Frans E, Kuja-Halkola R, Almqvist C, Sjolander A, Larsson H, Lichtenstein P. Paternal age at childbearing and offspring psychiatric and academic morbidity. JAMA Psychiatry 2014;71:432-438.
Dalman C, Allebeck P. Paternal age and schizophrenia: further support for an association. Am J Psychiatry 2002; 159:I591-I 592.
Daraki V, Roumeliotaki T, Koutra K, Georgiou V, Kampouri M, Kyriklaki A, Vafeiadi M, Papavasiliou S, Kogevinas M, Chatzi L. Effect of parental obesity and gestational diabetes on child neuropsychological and behavioral development at 4 years of age: the Rhea mother-child cohort, Crete, Greece. Eur Child Adolesc Psychiatry 2017;26:703-714.

Davey Smith G, Steer C, Leary S, Ness A. Is there an intrauterine influence on obesity? Evidence from parent child associations in the Avon Longitudinal Study of Parents and Children (ALSPAC). Arch Dis Child 2007;92:876-880.
de Jonge LL, Harris HR, Rich-Edwards JW, Willett WC, Forman MR, Jaddoe VW, Michels KB. Parental smoking in pregnancy and the risks of adult-onset hypertension. Hypertension 2013;61:494-500.
de Michelena MI, Burstein E, Lama JR, Vasquez JC. Paternal age as a risk factor for Down syndrome. Am J Med Genet 1993;45:679-682.
De Souza E, Alberman E, Morris JK. Down syndrome and paternal age, a new analysis of case-control data collected in the 1960s. Am J Med Genet A 2009; 149a: 1205-1208.
De Souza E, Morris JK. Case-control analysis of paternal age and trisomic anomalies. Arch Dis Child 2010;95:893-897.
Deng K, Liu Z, Lin Y, Mu D, Chen X, Li J, Li N, Deng Y, Li X, Wang Y et al. Periconceptional paternal smoking and the risk of congenital heart defects: a case-control study. Birth Defects Res A Clin Mol Teratol 2013;97:2I0-216.
DerKinderen DJ, Koten JW, Tan KE, Beemer FA, Van Romunde LK, Den Otter W. Parental age in sporadic hereditary retinoblastoma. Am J Ophthalmol 1990;110: 605-609.
Dockerty JD, Draper G, Vincent T, Rowan SD, Bunch KJ. Case-control study of parental age, parity and socioeconomic level in relation to childhood cancers. Int J Epidemiol 2001;30:1428-1437.
Durkin MS, Maenner MJ, Newschaffer CJ, Lee LC, Cunniff CM, Daniels JL, Kirby RS, Leavitt L, Miller L, Zahorodny W et al. Advanced parental age and the risk of autism spectrum disorder. Am J Epidemiol 2008;168:I268-1276.
Durmus B, Kruithof CJ, Gillman MH, Willemsen SP, Hofman A, Raat H, Eilers PH, Steegers EA, Jaddoe VW. Parental smoking during pregnancy, early growth, and risk of obesity in preschool children: the Generation R Study. Am J Clin Nutr 2011;94:164-171.
Durmus B, Arends LR, Ay L, Hokken-Koelega AC, Raat H, Hofman A, Steegers EA, Jaddoe VW. Parental anthropometrics, early growth and the risk of overweight in pre-school children: the Generation R Study. Pediatr Obes 2013;8: 339-350.
Dzurova D, Pikhart H. Down syndrome, paternal age and education: comparison of California and the Czech Republic. BMC Public Health 2005;5:69.
Ek M, Wicks S, Svensson AC, Idring S, Dalman C. Advancing paternal age and schizophrenia: the impact of delayed fatherhood. Schizophr Bull 2015;41: 708-714.
El-Saadi O, Pedersen CB, McNeil TF, Saha S, Welham J, O’Callaghan E, CantorGraae E, Chant D, Mortensen PB, McGrath J. Paternal and maternal age as risk factors for psychosis: findings from Denmark, Sweden and Australia. Schizophr Res 2004;67:227-236.
Erickson JD. Down syndrome, paternal age, maternal age and birth order. Ann Hum Genet I978;41:289-298.
Erickson JD. Paternal age and Down syndrome. Am J Hum Genet 1979;3 I:489-497. Erickson JD, Cohen MM Jr. A study of parental age effects on the occurrence of fresh mutations for the Apert syndrome. Ann Hum Genet 1974;38:89-96.
Erickson JD, Bjerkedal TO. Down syndrome associated with father's age in Norway. J Med Genet 198I; 18:22-28.
Eriksen W, Sundet JM, Tambs K. Paternal age at birth and the risk of obesity in young adulthood: a register-based birth cohort study of Norwegian males. Am J Hum Biol 2013;25:29-34.
Farioli A, Legittimo P, Mattioli S, Miligi L, Benvenuti A, Ranucci A, Salvan A, Rondelli R, Conter V, Magnani C. Tobacco smoke and risk of childhood acute lymphoblastic leukemia: findings from the SETIL case-control study. Cancer Causes Control 2014;25:683-692.
Figueiredo JC, Ly S, Magee KS, Ihenacho U, Baurley JW, Sanchez-Lara PA, Brindopke F, Nguyen TH, Nguyen V, Tangco MI et al. Parental risk factors for oral clefts among Central Africans, Southeast Asians, and Central Americans. Birth Defects Res A Clin Mol Teratol 2015; 103:863-879.
Filippini G, Maisonneuve P, McCredie M, Peris-Bonet R, Modan B, Preston-Martin S, Mueller BA, Holly EA, Cordier S, Choi NW et al. Relation of childhood brain tumors to exposure of parents and children to tobacco smoke: the SEARCH international case-control study. Surveillance of Environmental Aspects Related to Cancer in Humans. Int J Cancer 2002; 100:206-213.
Finley WH, Callahan A, Thompson JN. Parental age in the blepharophimosis, ptosis, epicanthus inversus, telecanthus complex. Am J Med Genet 1990;36: 414-417.

Fisch H, Hyun G, Golden R, Hensle TW, Olsson CA, Liberson GL. The influence of paternal age on down syndrome. J Urol 2003; 169:2275-2278.
Florath I, Kohler M, Weck MN, Brandt S, Rothenbacher D, Schottker B, Moss A, Gottmann P, Wabitsch M, Brenner H. Association of pre- and post-natal parental smoking with offspring body mass index: An 8-year follow-up of a birth cohort. Pediatr Obes 2014;9:|2|-134.
Foutz J, Mezuk B. Advanced paternal age and risk of psychotic-like symptoms in adult offspring. Schizophr Res 2015;165:123-127.
Frans EM, Sandin S, Reichenberg A, Lichtenstein P, Langstrom N, Hultman CM. Advancing paternal age and bipolar disorder. Arch Gen Psychiatry 2008;65: 1034-1040.
Frans EM, McGrath JJ, Sandin S, Lichtenstein P, Reichenberg A, Langstrom N, Hultman CM. Advanced paternal and grandpaternal age and schizophrenia: a three-generation perspective. Schizophr Res 201I;133:I20-124.
Frans EM, Sandin S, Reichenberg A, Langstrom N, Lichtenstein P, McGrath JJ, Hultman CM. Autism risk across generations: a population-based study of advancing grandpaternal and paternal age. JAMA Psychiatry 2013;70:516-521.
Frans E, MacCabe JH, Reichenberg A. Advancing paternal age and psychiatric disorders. World Psychiatry 2015;14:91-93.
Gaizauskiene A, Padaiga Z, Starkuviene S, Mizeriene R. Prediction of perinatal mortality at an early stage of pregnancy. Scand J Public Health 2007;35:564-569.
Gillberg C. Parental age in child psychiatric clinic attenders. Acta Psychiatr Scand 1982;66:471-478.
Gold EB, Leviton A, Lopez R, Gilles FH, Hedley-Whyte ET, Kolonel LN, Lyon JL, Swanson GM, Weiss NS, West D et al. Parental smoking and risk of childhood brain tumors. Am J Epidemiol 1993;137:620-628.
Goriely A, Wilkie AOM. Paternal age effect mutations and selfish spermatogonial selection: causes and consequences for human disease. Am J Hum Genet 2012; 90:175-200.
Green RF, Devine O, Crider KS, Olney RS, Archer N, Olshan AF, Shapira SK. Association of paternal age and risk for major congenital anomalies from the National Birth Defects Prevention Study, 1997 to 2004. Ann Epidemiol 2010;20: 24I-249.
Grether JK, Anderson MC, Croen LA, Smith D, Windham GC. Risk of autism and increasing maternal and paternal age in a large north American population. Am J Epidemiol 2009; 170: | I | 8-1| 26.
Grewal J, Carmichael SL, Yang W, Shaw GM. Paternal age and congenital malformations in offspring in California, 1989-2002. Matern Child Health J 2012;16:385-392.
Guyatt GH, Oxman AD, Vist GE, Kunz R, Falck-Ytter Y, Alonso-Coello P, Schunemann HJ. GRADE: an emerging consensus on rating quality of evidence and strength of recommendations. BMJ 2008;336:924-926.
Harville EW, Wilcox AJ, Lie RT, Abyholm F, Vindenes H. Epidemiology of cleft palate alone and cleft palate with accompanying defects. Eur J Epidemiol 2007;22: 389-395.
Heck JE, Lombardi CA, Meyers TJ, Cockburn M, Wilhelm M, Ritz B. Perinatal characteristics and retinoblastoma. Cancer Causes Control 2012;23:1567-I575.
Heppe DH, Kiefte-de Jong JC, Durmus B, Moll HA, Raat H, Hofman A, Jaddoe VW. Parental, fetal, and infant risk factors for preschool overweight: the Generation R Study. Pediatr Res 2013;73:120-127.
Herkrath AP, Herkrath FJ, Rebelo MA, Vettore MV. Parental age as a risk factor for non-syndromic oral clefts: a meta-analysis. J Dent 2012;40:3-14.
Hook EB, Cross PK, Lamson SH, Regal RR, Baird PA, Uh SH. Paternal age and Down syndrome in British Columbia. Am J Hum Genet 1981;33:123-128.
Hook EB, Cross PK. Paternal age and Down's syndrome genotypes diagnosed prenatally: no association in New York state data. Hum Genet 1982;62:167-174.
Hook EB, Regal RR. A search for a paternal-age effect upon cases of $47,+21$ in which the extra chromosome is of paternal origin. Am J Hum Genet 1984;36:413-42I.
Horta BL, Victora CG, Menezes AM, Halpern R, Barros FC. Low birthweight, preterm births and intrauterine growth retardation in relation to maternal smoking. Paediatr Perinat Epidemiol I997; II:I40-15I.
Howe LD, Matijasevich A, Tilling K, Brion MJ, Leary SD, Smith GD, Lawlor DA. Maternal smoking during pregnancy and offspring trajectories of height and adiposity: comparing maternal and paternal associations. Int J Epidemiol 2012;4I: 722-732.
Hu J, Mao Y, Ugnat AM. Parental cigarette smoking, hard liquor consumption and the risk of childhood brain tumors-a case-control study in northeast China. Acta Oncol 2000;39:979-984.

Hultman CM, Sandin S, Levine SZ, Lichtenstein P, Reichenberg A. Advancing paternal age and risk of autism: new evidence from a population-based study and a meta-analysis of epidemiological studies. Mol Psychiatry 2011;16:1203-1212.
Hvolgaard Mikkelsen S, Olsen J, Bech BH, Obel C. Parental age and attention-deficit/hyperactivity disorder (ADHD). Int J Epidemiol 2016;46:409-420.
Idring S, Magnusson C, Lundberg M, Ek M, Rai D, Svensson AC, Dalman C, Karlsson H, Lee BK. Parental age and the risk of autism spectrum disorders: findings from a Swedish population-based cohort. Int J Epidemiol 2014;43:I07-II5.
Inoue S, Naruse H, Yorifuji T, Kato T, Murakoshi T, Doi H, Subramanian SV. Impact of maternal and paternal smoking on birth outcomes. J Public Health (Oxf) 2016;39:1-10.
Iwayama M, Kira R, Kinukawa N, Sakai Y, Torisu H, Sanefuji M, Ishizaki Y, Nose Y, Matsumoto T, Hara T. Parental age and child growth and development: child health check-up data. Pediatr Int 2011;53:709-714.
Jaaskelainen A, Pussinen J, Nuutinen O, Schwab U, Pirkola J, Kolehmainen M, Jarvelin MR, Laitinen J. Intergenerational transmission of overweight among Finnish adolescents and their parents: a 16-year follow-up study. Int J Obes (Lond) 2011;35:1289-1294.
Jacobsson B, Ladfors L, Milsom I. Advanced maternal age and adverse perinatal outcome. Obstet Gynecol 2004; 104:727-733.
Javaras KN, Rickert ME, Thornton LM, Peat CM, Baker JH, Birgegard A, Norring C, Landen M, Almqvist C, Larsson H et al. Paternal age at childbirth and eating disorders in offspring. Psychol Med 2017;47:576-584.
Ji BT, Shu XO, Linet MS, Zheng W, Wacholder S, Gao YT, Ying DM, Jin F. Paternal cigarette smoking and the risk of childhood cancer among offspring of nonsmoking mothers. J Natl Cancer Inst 1997;89:238-244.
John EM, Savitz DA, Sandler DP. Prenatal exposure to parents’ smoking and childhood cancer. Am J Epidemiol I99।;133:|23-132.
Johnson KJ, Carozza SE, Chow EJ, Fox EE, Horel S, McLaughlin CC, Mueller BA, Puumala SE, Reynolds P, Von Behren J et al. Parental age and risk of childhood cancer: a pooled analysis. Epidemiol 2009;20:475-483.
Johnson KJ, Williams KS, Ross JA, Krailo MD, Tomlinson GE, Malogolowkin MH, Feusner JH, Spector LG. Parental tobacco and alcohol use and risk of hepatoblastoma in offspring: a report from the children's oncology group. Cancer Epidemiol, Biomarkers Prev 2013;22:1837-I843.
Kazaura MR, Lie RT. Down's syndrome and paternal age in Norway. Paediatr Perinat Epidemiol 2002;16:314-319.
Kazaura MR, Lie RT, Irgens LM, Didriksen A, Kapstad M, Egenaes J, Bjerkedal T. Increasing risk of gastroschisis in Norway: an age-period-cohort analysis. Am J Epidemiol 2004a; I59:358-363.
Kazaura M, Lie RT, Skjaerven R. Paternal age and the risk of birth defects in Norway. Ann Epidemiol 2004b; 14:566-570.
Khandwala YS, Zhang CA, Lu Y, Eisenberg ML. The age of fathers in the USA is rising: an analysis of 168867480 births from 1972 to 2015. Hum Reprod 2017;32: 2110-2116.
King MD, Fountain C, Dakhlallah D, Bearman PS. Estimated autism risk and older reproductive age. Am J Public Health 2009;99:1673-1679.
Klebanoff MA, Mednick BR, Schulsinger C, Secher NJ, Shiono PH. Father's effect on infant birth weight. Am J Obstet Gynecol 1998;178:I022-1026.
Ko TJ, Tsai LY, Chu LC, Yeh SJ, Leung C, Chen CY, Chou HC, Tsao PN, Chen PC, Hsieh WS. Parental smoking during pregnancy and its association with low birth weight, small for gestational age, and preterm birth offspring: a birth cohort study. Pediatr Neonatol 2014;55:20-27.
Kong A, Frigge ML, Masson G, Besenbacher S, Sulem P, Magnusson G, Gudjonsson SA, Sigurdsson A, Jonasdottir A, Jonasdottir A et al. Rate of de novo mutations and the importance of father's age to disease risk. Nature 2012;488:47I-475.
Krapels IP, Zielhuis GA, Vroom F, de Jong-van den Berg LT, Kuijpers-Jagtman AM, van der Molen AB, Steegers-Theunissen RP. Periconceptional health and lifestyle factors of both parents affect the risk of live-born children with orofacial clefts. Birth Defects Res A Clin Mol Teratol 2006;76:6I3-620.
Kuciene R, Dulskiene V. Parental cigarette smoking and the risk of congenital heart septal defects. Medicina (Kaunas) 2010;46:635-641.
Kwok MK, Schooling CM, Lam TH, Leung GM. Paternal smoking and childhood overweight: evidence from the Hong Kong 'Children of 1997'. Pediatrics 2010; 126: 46 -e56.
L'Abee C, Vrieze I, Kluck T, Erwich JJ, Stolk RP, Sauer PJ. Parental factors affecting the weights of the placenta and the offspring. J Perinat Med 201 I;39:27-34.

Lampi KM, Hinkka-Yli-Salomaki S, Lehti V, Helenius H, Gissler M, Brown AS, Sourander A. Parental age and risk of autism spectrum disorders in a Finnish national birth cohort. J Autism Dev Disord 20I3;43:2526-2535.
Langley K, Heron J, Smith GD, Thapar A. Maternal and paternal smoking during pregnancy and risk of ADHD symptoms in offspring: testing for intrauterine effects. Am J Epidemiol 2012;176:261-268.
Larfors G, Hallbook H, Simonsson B. Parental age, family size, and offspring's risk of childhood and adult acute leukemia. Cancer Epidemiol, Biomarkers Prev 2012; 21:1185-1190.
Larsson HJ, Eaton WW, Madsen KM, Vestergaard M, Olesen AV, Agerbo E, Schendel D, Thorsen P, Mortensen PB. Risk factors for autism: perinatal factors, parental psychiatric history, and socioeconomic status. Am J Epidemiol 2005; $\mathbf{1 6}$ I: 916-925; discussion 926-918.
Lauritsen MB, Pedersen CB, Mortensen PB. Effects of familial risk factors and place of birth on the risk of autism: a nationwide register-based study. J Child Psychol Psychiatry 2005;46:963-97I.
Lawlor DA, Smith GD, O'Callaghan M, Alati R, Mamun AA, Williams GM, Najman JM. Epidemiologic evidence for the fetal overnutrition hypothesis: findings from the mater-university study of pregnancy and its outcomes. Am J Epidemiol 2007; 165:418-424.
Lawlor DA, Timpson NJ, Harbord RM, Leary S, Ness A, McCarthy MI, Frayling TM, Hattersley AT, Smith GD. Exploring the developmental overnutrition hypothesis using parental-offspring associations and FTO as an instrumental variable. PLoS Med 2008;5:e33.
Leary SD, Smith GD, Rogers IS, Reilly JJ, Wells JC, Ness AR. Smoking during pregnancy and offspring fat and lean mass in childhood. Obesity (Silver Spring) 2006; 14:2284-2293.
Lee KM, Ward MH, Han S, Ahn HS, Kang HJ, Choi HS, Shin HY, Koo HH, Seo JJ, Choi JE et al. Paternal smoking, genetic polymorphisms in CYPIAI and childhood leukemia risk. Leuk Res 2009;33:250-258.
Lehrer DS, Pato MT, Nahhas RW, Miller BR, Malaspina D, Buckley PF, Sobell JL, Walsh-Messinger J, Cohort Consortium GP, Pato CN. Paternal age effect: replication in schizophrenia with intriguing dissociation between bipolar with and without psychosis. Am J Med Genet B Neuropsychiatr Genet 2016; I7 I:495-505.
Lian ZH, Zack MM, Erickson JD. Paternal age and the occurrence of birth defects. Am J Hum Genet 1986;39:648-660.
Linabery AM, Nahhas RW, Johnson W, Choh AC, Towne B, Odegaard AO, Czerwinski SA, Demerath EW. Stronger influence of maternal than paternal obesity on infant and early childhood body mass index: the Fels Longitudinal Study. Pediatr Obes 2013;8:159-169.
Liu R, Zhang L, McHale CM, Hammond SK. Paternal smoking and risk of childhood acute lymphoblastic leukemia: systematic review and meta-analysis. J Oncol 2011; 2011:854584.
Lorda-Sanchez I, Prieto L, Rodriguez-Pinilla E, Martinez-Frias ML. Increased parental age and number of pregnancies in Klippel-Trenaunay-Weber syndrome. Ann Hum Genet 1998;62:235-239.
Lowe X, Eskenazi B, Nelson DO, Kidd S, Alme A, Wyrobek AJ. Frequency of XY sperm increases with age in fathers of boys with Klinefelter syndrome. Am J Hum Genet 2001;69:1046-1054.
Lundstrom S, Haworth CM, Carlstrom E, Gillberg C, Mill J, Rastam M, Hultman CM, Ronald A, Anckarsater H, Plomin R et al. Trajectories leading to autism spectrum disorders are affected by paternal age: findings from two nationally representative twin studies. J Child Psychol Psychiatry 20I0;51:850-856.
MacArthur AC, McBride ML, Spinelli JJ, Tamaro S, Gallagher RP, Theriault G. Risk of childhood leukemia associated with parental smoking and alcohol consumption prior to conception and during pregnancy: the cross-Canada childhood leukemia study. Cancer Causes Control 2008; 19:283-295.
Magnani C, Pastore G, Luzzatto L, Terracini B. Parental occupation and other environmental factors in the etiology of leukemias and non-Hodgkin's lymphomas in childhood: a case-control study. Tumori 1990;76:413-419.
Magnus P, Berg K, Bjerkedal T, Nance WE. Parental determinants of birth weight. Clin Genet 1984;26:397-405.
Maimburg RD, Vaeth M. Perinatal risk factors and infantile autism. Acta Psychiatr Scand 2006; I I4:257-264.
Malaspina D, Harlap S, Fennig S, Heiman D, Nahon D, Feldman D, Susser ES. Advancing paternal age and the risk of schizophrenia. Arch Gen Psychiatry 200I; 58:36I-367.

Mamidala MP, Polinedi A, P T V PK, Rajesh N, Vallamkonda OR, Udani V, Singhal N, Rajesh V. Prenatal, perinatal and neonatal risk factors of Autism Spectrum Disorder: a comprehensive epidemiological assessment from India. Res Dev Disabil 20|3;34:3004-30|3.
Martinez FD, Wright AL, Taussig LM. The effect of paternal smoking on the birthweight of newborns whose mothers did not smoke. Group Health Medical Associates. Am J Public Health 1994;84:I489-I49।.
Materna-Kiryluk A, Wisniewska K, Badura-Stronka M, Mejnartowicz J, Wieckowska B, Balcar-Boron A, Czerwionka-Szaflarska M, Gajewska E, Godula-Stuglik U, Krawczynski M et al. Parental age as a risk factor for isolated congenital malformations in a Polish population. Paediatr Perinat Epidemiol 2009;23:29-40.
Matsunaga E, Minoda K, Sasaki MS. Parental age and seasonal variation in the births of children with sporadic retinoblastoma: a mutation-epidemiologic study. Hum Genet 1990;84:155-158.
Mattioli S, Farioli A, Legittimo P, Miligi L, Benvenuti A, Ranucci A, Salvan A, Rondelli R, Magnani C. Tobacco smoke and risk of childhood acute nonlymphocytic leukemia: findings from the SETIL study. PLoS One 2014;9: elllo28.
Maule MM, Vizzini L, Merletti F, Magnani C, Pastore G, Richiardi L. Parental age and risk of acute lymphocytic leukaemia and embryonal tumours in the Piedmont Region, Italy. Int J Epidemiol 2007;36:691-692.
McCredie M, Maisonneuve P, Boyle P. Antenatal risk factors for malignant brain tumours in New South Wales children. Int J Cancer 1994;56:6-10.
McGrath JJ, Petersen L, Agerbo E, Mors O, Mortensen PB, Pedersen CB. A comprehensive assessment of parental age and psychiatric disorders. JAMA Psychiatry 2014;71:301-309.
McIntosh GC, Olshan AF, Baird PA. Paternal age and the risk of birth defects in offspring. Epidemiol 1995;6:282-288.
Menegaux F, Ripert M, Hemon D, Clavel J. Maternal alcohol and coffee drinking, parental smoking and childhood leukaemia: a French population-based case-control study. Paediatr Perinat Epidemiol 2007;21:293-299.
Menezes PR, Lewis G, Rasmussen F, Zammit S, Sipos A, Harrison GL, Tynelius P, Gunnell D. Paternal and maternal ages at conception and risk of bipolar affective disorder in their offspring. Psychol Med 2010;40:477-485.
Metayer C, Zhang L, Wiemels JL, Bartley K, Schiffman J, Ma X, Aldrich MC, Chang JS, Selvin S, Fu CH et al. Tobacco smoke exposure and the risk of childhood acute lymphoblastic and myeloid leukemias by cytogenetic subtype. Cancer Epidemiol, Biomarkers Prev 2013;22:1600-1611.
Metayer C, Petridou E, Arangure JM, Roman E, Schuz J, Magnani C, Mora AM, Mueller BA, de Oliveira MS, Dockerty JD et al. Parental tobacco smoking and acute myeloid leukemia: the Childhood Leukemia International Consortium. Am J Epidemiol 2016;184:26I-273.
Miller B, Alaraisanen A, Miettunen J, Jarvelin MR, Koponen H, Rasanen P, Isohanni M, Kirkpatrick B. Advanced paternal age, mortality, and suicide in the general population. J Nerv Ment Dis 2010;198:404-4II.
Miller B, Messias E, Miettunen J, Alaraisanen A, Jarvelin MR, Koponen H, Rasanen P, Isohanni M, Kirkpatrick B. Meta-analysis of paternal age and schizophrenia risk in male versus female offspring. Schizophr Bull 2011;37:1039-1047.
Milne E, Greenop KR, Scott RJ, Bailey HD, Attia J, Dalla-Pozza L, de Klerk NH, Armstrong BK. Parental prenatal smoking and risk of childhood acute lymphoblastic leukemia. Am J Epidemiol 2012; 175:43-53.
Milne E, Greenop KR, Scott RJ, Ashton LJ, Cohn RJ, de Klerk NH, Armstrong BK. Parental smoking and risk of childhood brain tumors. Int J Cancer 2013;133: 253-259.
Mok PL, Antonsen S, Pedersen CB, Webb RT. Younger or older parental age and risk of suicidality, premature death, psychiatric illness, and criminality in offspring. J Affect Disord 2017;208:I30-138.
Morrison J, Williams GM, Najman JM, Andersen MJ. The influence of paternal height and weight on birth-weight. Aust N Z J Obstet Gynaecol 1991;3 I:II4-116.
Mutsaerts MA, Groen H, Buiter-Van der Meer A, Sijtsma A, Sauer PJ, Land JA, Mol BW, Corpeleijn E, Hoek A. Effects of paternal and maternal lifestyle factors on pregnancy complications and perinatal outcome. A population-based birthcohort study: the GECKO Drenthe cohort. Hum Reprod 2014;29:824-834.
Nahum GG, Stanislaw H. Relationship of paternal factors to birth weight. J Reprod Med 2003;48:963-968.
Naserbakht M, Ahmadkhaniha HR, Mokri B, Smith CL. Advanced paternal age is a risk factor for schizophrenia in Iranians. Ann Gen Psychiatry 2011;10:15.

National Institute for Health and Welfare. Nordic perinatal statistics 2014. Helsinki, Finland. https://www.thl.fi/en/web/thlfi-en/statistics/statistics-by-topic/sexual-and-reproductive-health/parturients-deliveries-and-births/nordic-perinatal-statistics.
Nilsen AB, Waldenstrom U, Rasmussen S, Hjelmstedt A, Schytt E. Characteristics of first-time fathers of advanced age: a Norwegian population-based study. BMC Pregnancy Childbirth 2013;13:29.
Nomura Y, Marks DJ, Halperin JM. Prenatal exposure to maternal and paternal smoking on attention deficit hyperactivity disorders symptoms and diagnosis in offspring. J Nerv Ment Dis 2010; 198:672-678.
Norman MA, Holly EA, Ahn DK, Preston-Martin S, Mueller BA, Bracci PM. Prenatal exposure to tobacco smoke and childhood brain tumors: results from the United States West Coast childhood brain tumor study. Cancer Epidemiol, Biomarkers Prev 1996;5:127-133.
Nybo Andersen AM, Hansen KD, Andersen PK, Davey Smith G. Advanced paternal age and risk of fetal death: a cohort study. Am J Epidemiol 2004;160:1214-1222.
Nybo Andersen AM, Urhoj SK. Is advanced paternal age a health risk for the offspring? Fertil Steril 2017;107:3I2-3I8.
O'Callaghan MJ, Williams GM, Andersen MJ, Bor W, Najman JM. Prediction of obesity in children at 5 years: a cohort study. J Paediatr Child Health 1997;33: 3|I-316.
Olshan AF, Schnitzer PG, Baird PA. Paternal age and the risk of congenital heart defects. Teratology 1994;50:80-84.
Olshan AF, Ananth CV, Savitz DA. Intrauterine growth retardation as an endpoint in mutation epidemiology: an evaluation based on paternal age. Mutat Res 1995; 344:89-94.
Orioli IM, Castilla EE, Scarano G, Mastroiacovo P. Effect of paternal age in achondroplasia, thanatophoric dysplasia, and osteogenesis imperfecta. Am J Med Genet 1995;59:209-217.
Orsi L, Rudant J, Ajrouche R, Leverger G, Baruchel A, Nelken B, Pasquet M, Michel G, Bertrand Y, Ducassou S et al. Parental smoking, maternal alcohol, coffee and tea consumption during pregnancy, and childhood acute leukemia: the ESTELLE study. Cancer Causes Control 2015;26:I003-1017.
Pang D, McNally R, Birch JM. Parental smoking and childhood cancer: results from the United Kingdom Childhood Cancer Study. Br J Cancer 2003;88:373-38।.
Parner ET, Baron-Cohen S, Lauritsen MB, Jorgensen M, Schieve LA, YearginAllsopp M, Obel C. Parental age and autism spectrum disorders. Ann Epidemiol 2012;22:143-150.
Patro B, Liber A, Zalewski B, Poston L, Szajewska H, Koletzko B. Maternal and paternal body mass index and offspring obesity: a systematic review. Ann Nutr Metab 20I3;63:32-4I.
Perrin MC, Brown AS, Malaspina D. Aberrant epigenetic regulation could explain the relationship of paternal age to schizophrenia. Schizophr Bull 2007; 33:1270-1273.
Petersen L, Mortensen PB, Pedersen CB. Paternal age at birth of first child and risk of schizophrenia. Am J Psychiatry 201 I; 168:82-88.
Plichart M, Menegaux F, Lacour B, Hartmann O, Frappaz D, Doz F, Bertozzi AI, Defaschelles AS, Pierre-Kahn A, Icher C et al. Parental smoking, maternal alcohol, coffee and tea consumption during pregnancy and childhood malignant central nervous system tumours: the ESCALE study (SFCE). Eur J Cancer Prev 2008; 17:376-383.
Polednak AP. Paternal age in relation to selected birth defects. Hum Biol 1976;48: 727-739.
Poletta FA, Castilla EE, Orioli IM, Lopez-Camelo JS. Regional analysis on the occurrence of oral clefts in South America. Am J Med Genet A 2007; I43a:32 I6-3227.
Pritchard CW, Sutherland HW, Carr-Hill RA. Birthweight and paternal height. Br J Obstet Gynaecol 1983;90:156-161.
Quinlan CA, McVeigh KH, Driver CR, Govind P, Karpati A. Parental Age and Autism Spectrum Disorders Among New York City Children 0-36 Months of Age. Matern Child Health J 2015;19:1783-1790.
Racine SE, Culbert KM, Burt SA, Klump KL. Advanced paternal age at birth: phenotypic and etiologic associations with eating pathology in offspring. Psychol Med 2014;44:1029-1041.
Reichenberg A, Gross R, Weiser M, Bresnahan M, Silverman J, Harlap S, Rabinowitz J, Shulman C, Malaspina D, Lubin G et al. Advancing paternal age and autism. Arch Gen Psychiatry 2006;63: I026-1032.
Reichenberg A, Gross R, Sandin S, Susser ES. Advancing paternal and maternal age are both important for autism risk. Am J Public Health 2010;100:772-773.

Reichman NE, Teitler JO. Paternal age as a risk factor for low birthweight. Am J Public Health 2006;96:862-866.
Reilly JJ, Armstrong J, Dorosty AR, Emmett PM, Ness A, Rogers I, Steer C, Sherriff A. Early life risk factors for obesity in childhood: cohort study. BMJ 2005;330:I357.
Riccardi VM, Dobson ICE, Chakraborty R, Bontke C. The pathophysiology of neurofibromatosis: IX. Paternal age as a factor in the origin of new mutations. Am J Med Genet 1984; 18:169-176.
Roecker GO, Huether CA. An analysis for paternal-age effect in Ohio's Down syndrome births, 1970-1980. Am J Hum Genet 1983;35: I297-1306.
Roth MP, Stoll C, Taillemite JL, Girard S, Boue A. Paternal age and Down's syndrome diagnosed prenatally: no association in French data. Prenat Diagn 1983a; 3:327-335.
Roth MP, Feingold J, Baumgarten A, Bigel P, Stoll C. Reexamination of paternal age effect in Down's syndrome. Hum Genet 1983b;63:149-152.
Rudant J, Menegaux F, Leverger G, Baruchel A, Lambilliotte A, Bertrand Y, Patte C, Pacquement H, Verite C, Robert A et al. Childhood hematopoietic malignancies and parental use of tobacco and alcohol: the ESCALE study (SFCE). Cancer Causes Control 2008;19:1277-1290.
Saha S, Barnett AG, Foldi C, Burne TH, Eyles DW, Buka SL, McGrath JJ. Advanced paternal age is associated with impaired neurocognitive outcomes during infancy and childhood. PLoS Med 2009;6:e40.
Sandin S, Schendel D, Magnusson P, Hultman C, Suren P, Susser E, Gronborg T, Gissler M, Gunnes N, Gross R et al. Autism risk associated with parental age and with increasing difference in age between the parents. Mol Psychiatry 2016; 21:693-700.
Sartorius GA, Nieschlag E. Paternal age and reproduction. Hum Reprod Update 2010;16:65-79.
Sasanfar R, Haddad SA, Tolouei A, Ghadami M, Yu D, Santangelo SL. Paternal age increases the risk for autism in an Iranian population sample. Mol Autism 2010;1:2.
Savitz DA, Schwingl PJ, Keels MA. Influence of paternal age, smoking, and alcohol consumption on congenital anomalies. Teratology 1991;44:429-440.
Schuz J, Kaatsch P, Kaletsch U, Meinert R, Michaelis J. Association of childhood cancer with factors related to pregnancy and birth. Int J Epidemiol 1999;28:63I-639.
Selvin S, Garfinkel J. The relationship between parental age and birth order with the percentage of low birth-weight infants. Hum Biol 1972;44:50I-509.
Sergentanis TN, Thomopoulos TP, Gialamas SP, Karalexi MA, Biniaris-Georgallis SI, Kontogeorgi E, Papathoma P, Tsilimidos G, Skalkidou A, lliadou AN et al. Risk for childhood leukemia associated with maternal and paternal age. Eur J Epidemiol 2015;30:1229-1261.
Shah PS. Paternal factors and low birthweight, preterm, and small for gestational age births: a systematic review. Am J Obstet Gynecol 2010;202:103-I23.
Shelton JF, Tancredi DJ, Hertz-Picciotto I. Independent and dependent contributions of advanced maternal and paternal ages to autism risk. Autism Res 2010;3:30-39.
Shu XO, Ross JA, Pendergrass TW, Reaman GH, Lampkin B, Robison LL. Parental alcohol consumption, cigarette smoking, and risk of infant leukemia: a Childrens Cancer Group study. J Natl Cancer Inst 1996;88:24-3I.
Sinclair KD, Watkins AJ. Parental diet, pregnancy outcomes and offspring health: metabolic determinants in developing oocytes and embryos. Reprod Fertil Dev 2013;26:99-114.
Sipos A, Rasmussen F, Harrison G, Tynelius P, Lewis G, Leon DA, Gunnell D. Paternal age and schizophrenia: a population based cohort study. BMJ 2004;329: 1070.

Sorahan T, Lancashire RJ, Hulten MA, Peck I, Stewart AM. Childhood cancer and parental use of tobacco: deaths from 1953 to 1955. BrJ Cancer 1997a; 75:134-138.
Sorahan T, Prior P, Lancashire RJ, Faux SP, Hulten MA, Peck IM, Stewart AM. Childhood cancer and parental use of tobacco: deaths from 1971 to 1976. Br J Cancer 1997b;76:1525-I53I.
Sorahan T, McKinney PA, Mann JR, Lancashire RJ, Stiller CA, Birch JM, Dodd HE, Cartwright RA. Childhood cancer and parental use of tobacco: findings from the inter-regional epidemiological study of childhood cancer (IRESCC). Br J Cancer 2001;84:14I-146.
Sorahan T, Lancashire RJ. Parental cigarette smoking and childhood risks of hepatoblastoma: OSCC data. BrJ Cancer 2004;90:1016-1018.

Sorensen HJ, Pedersen CB, Nordentoft M, Mortensen PB, Ehrenstein V, Petersen L. Effects of paternal age and offspring cognitive ability in early adulthood on the risk of schizophrenia and related disorders. Schizophr Res 2014;160:131-135.
Soubry A, Verbeke G, Hoyo C. Do early paternal exposures to lifestyle factors such as smoking increase the risk of chronic diseases in the offspring? Eur J Hum Genet 2014;22:134I-1342.
Steegers-Theunissen RP, Twigt J, Pestinger V, Sinclair KD. The periconceptional period, reproduction and long-term health of offspring: the importance of onecarbon metabolism. Hum Reprod Update 2013; 19:640-655.
Stene J, Fischer G, Stene E, Mikkelsen M, Petersen E. Paternal age effect in Down's syndrome. Ann Hum Genet 1977;40:299-306.
Stene J, Stene E, Stengel-Rutkowski S, Murken JD. Paternal age and Down's syndrome: data from prenatal diagnoses (DFG). Hum Genet 198।;59:I 19-124.
Stene LC, Magnus P, Lie RT, Sovik O, Joner G. Maternal and paternal age at delivery, birth order, and risk of childhood onset type I diabetes: population based cohort study. BMJ 2001;323:369.
Stern JE, Luke B, Hornstein MD, Cabral H, Gopal D, Diop H, Kotelchuck M. The effect of father's age in fertile, subfertile, and assisted reproductive technology pregnancies: a population based cohort study. J Assist Reprod Genet 2014;3I: 1437-1444.
Stroup DF, Berlin JA, Morton SC, Olkin I, Williamson GD, Rennie D, Moher D, Becker BJ, Sipe TA, Thacker SB. Meta-analysis of observational studies in epidemiology: a proposal for reporting. Meta-analysis Of Observational Studies in Epidemiology (MOOSE) group. JAMA 2000;283:2008-20I2.
Su XJ, Yuan W, Huang GY, Olsen J, Li J. Paternal age and offspring congenital heart defects: a national cohort study. PLoS One 2015;10:e0121030.
Suren P, Gunnes N, Roth C, Bresnahan M, Hornig M, Hirtz D, Lie KK, Lipkin WI, Magnus P, Reichborn-Kjennerud T et al. Parental obesity and risk of autism spectrum disorder. Pediatrics 2014;133:ell28-ell38.
Taal HR, de Jonge LL, van Osch-Gevers L, Steegers EA, Hofman A, Helbing WA, van der Heijden AJ, Jaddoe VW. Parental smoking during pregnancy and cardiovascular structures and function in childhood: the Generation R Study. Int J Epidemiol 2013;42:1371-1380.
Takano T, Kawashima T, Yamanouchi Y, Kitayama K, Baba T, Ueno K, Hamaguchi H. Genetics of neurofibromatosis I in Japan: mutation rate and paternal age effect. Hum Genet 1992;89:28I-286.
Tang ML. Environmental tobacco smoke and child development: a case-control study on Hong Kong Chinese toddlers. Hong Kong J Paediatr 2006; I I; II8-127+160-161.
Tay JS, Yip WC, Joseph R. Parental age and birth order in Chinese children with congenital heart disease. J Med Genet I982; 19:44I-443.
Tellier AL, Lyonnet S, Cormier-Daire V, de Lonlay P, Abadie V, Baumann C, Bonneau D, Labrune P, Lacombe D, Le Merrer M et al. Increased paternal age in CHARGE association. Clin Genet 1996;50:548-550.
Teras LR, Gaudet MM, Blase JL, Gapstur SM. Parental age at birth and risk of hematological malignancies in older adults. Am J Epidemiol 2015;182:41-48.
Tiesler CM, Chen CM, Sausenthaler S, Herbarth O, Lehmann I, Schaaf B, Kramer U, von Berg A, von Kries R, Wichmann HE et al. Passive smoking and behavioural problems in children: results from the LISAplus prospective birth cohort study. Environ Res 2011; I I I:II73-1I79.
To WW, Cheung W, Kwok JS. Paternal height and weight as determinants of birth weight in a Chinese population. Am J Perinatol 1998; 15:545-548.
Torrey EF, Buka S, Cannon TD, Goldstein JM, Seidman LJ, Liu T, Hadley T, Rosso IM, Bearden C, Yolken RH. Paternal age as a risk factor for schizophrenia: how important is it? Schizophr Res 2009; I I4:I-5.
Toschke AM, Ehlin A, Koletzko B, Montgomery SM. Paternal smoking is associated with a decreased prevalence of type I diabetes mellitus among offspring in two national British birth cohort studies (NCDS and BCS70). J Perinat Med 2007;35: 43-47.
Tough SC, Faber AJ, Svenson LW, Johnston DW. Is paternal age associated with an increased risk of low birthweight, preterm delivery, and multiple birth? Can J Public Health 2003;94:88-92.
Tsuchiya KJ, Takagai S, Kawai M, Matsumoto H, Nakamura K, Minabe Y, Mori N, Takei N. Advanced paternal age associated with an elevated risk for schizophrenia in offspring in a Japanese population. Schizophr Res 2005;76:337-342.
Tsuchiya KJ, Matsumoto K, Miyachi T, Tsujii M, Nakamura K, Takagai S, Kawai M, Yagi A, Iwaki K, Suda S et al. Paternal age at birth and high-functioning autisticspectrum disorder in offspring. BrJ Psychiatry 2008;193:316-321.

Urhoj SK, Jespersen LN, Nissen M, Mortensen LH, Nybo Andersen AM. Advanced paternal age and mortality of offspring under 5 years of age: a register-based cohort study. Hum Reprod 2014;29:343-350.
Urhoj SK, Mortensen LH, Nybo Andersen AM. Advanced paternal age and risk of musculoskeletal congenital anomalies in offspring. Birth Defects Res B Dev Reprod Toxicol 2015; 104:273-280.
Urhoj SK, Andersen PK, Mortensen LH, Davey Smith G, Nybo Andersen AM. Advanced paternal age and stillbirth rate: a nationwide register-based cohort study of 944,03I pregnancies in Denmark. Eur J Epidemiol 2017a;32:227-234.
Urhoj SK, Raaschou-Nielsen O, Hansen AV, Mortensen LH, Andersen PK, Nybo Andersen AM. Advanced paternal age and childhood cancer in offspring: A nationwide register-based cohort study. Int J Cancer 2017b; 140:246I-2472.
van Balkom ID, Bresnahan M, Vuijk PJ, Hubert J, Susser E, Hoek HW. Paternal age and risk of autism in an ethnically diverse, non-industrialized setting: Aruba. PLoS One 2012;7:e45090.
van Rooij IA, Wijers CH, Rieu PN, Hendriks HS, Brouwers MM, Knoers NV, de Blaauw I, Roeleveld N. Maternal and paternal risk factors for anorectal malformations: a Dutch case-control study. Birth Defects Res A Clin Mol Teratol 2010; 88:152-158.
Vashist M, Neelkamal, Kumar A. Whether paternal age effect exists as a risk factor in trisomy 21. Biosci Biotechnol Res Asia 2011;8:247-252.
Wang SH, Liu CM, Hwu HG, Hsiao CK, Chen WJ. Association of older paternal age with earlier onset among co-affected schizophrenia sib-pairs. Psychol Med 2015;45:2205-2213.
Wasserman CR, Shaw GM, O'Malley CD, Tolarova MM, Lammer EJ. Parental cigarette smoking and risk for congenital anomalies of the heart, neural tube, or limb. Teratology 1996;53:26I-267.
Waye MMY, Cheng HY. Genetics and epigenetics of autism (Review). Psychiatry Clin Neurosci 2017.
Wennberg AL, Opdahl S, Bergh C, Aaris Henningsen AK, Gissler M, Romundstad LB, Pinborg A, Tiitinen A, Skjaerven R, Wennerholm UB. Effect of maternal age on maternal and neonatal outcomes after assisted reproductive technology. Fertil Steril 2016; 106: | 142 - I | 49. e I I I4.
Wilcox MA, Newton CS, Johnson IR. Paternal influences on birthweight. Acta Obstet Gynecol Scand 1995;74:15-18.
Winikoff B, Debrovner CH. Anthropometric determinants of birth weight. Obstet Gynecol 198।;58:678-684.
Wohl M, Gorwood P. Paternal ages below or above 35 years old are associated with a different risk of schizophrenia in the offspring. Eur Psychiatry 2007;22:22-26.
Wolf CM. Paternal age effect for cleft lip and palate. Am J Hum Genet 1963;15: 389-393.
Wu Y, Liu X, Luo H, Deng W, Zhao G, Wang Q, Zhang L, Ma X, Liu X, Murray RA et al. Advanced paternal age increases the risk of schizophrenia and obsessive-compulsive disorder in a Chinese Han population. Psychiatry Res 2012; 198:353-359.
Wu S, Wu F, Ding Y, Hou J, Bi J, Zhang Z. Advanced parental age and autism risk in children: a systematic review and meta-analysis. Acta Psychiatr Scand 2017; 135:29-41.
Wunsch G, Gourbin C. Parents' age at birth of their offspring and child survival. Soc Biol 2002;49:174-184.
Yang Q, Olshan AF, Bondy ML, Shah NR, Pollock BH, Seeger RC, Look AT, Cohn SL. Parental smoking and alcohol consumption and risk of neuroblastoma. Cancer Epidemiol, Biomarkers Prev 2000;9:967-972.
Yang Q, Wen SW, Leader A, Chen XK, Lipson J, Walker M. Paternal age and birth defects: how strong is the association? Hum Reprod 2007;22:696-70I.
Yeung EH, Sundaram R, Ghassabian A, Xie Y, Buck Louis G. Parental obesity and early childhood development. Pediatrics 2017;139:1-12.
Yip BH, Pawitan Y, Czene K. Parental age and risk of childhood cancers: a population-based cohort study from Sweden. Int J Epidemiol 2006;35: 1495-1503.
Zakar R, Zakar MZ, Aqil N, Nasrullah M. Paternal factors associated with neonatal deaths and births with low weight: evidence from Pakistan Demographic and Health Survey 2006-2007. Matern Child Health J 2015; 19:1634-1642.
Zammit S, Allebeck P, Dalman C, Lundberg I, Hemmingson T, Owen MJ, Lewis G. Paternal age and risk for schizophrenia. BrJ Psychiatry 2003; I83:405-408.
Zenzes MT. Smoking and reproduction: gene damage to human gametes and embryos. Hum Reprod Update 2000;6:I22-I31.

Zhan SY, Lian ZH, Zheng DZ, Gao L. Effect of fathers' age and birth order on occurrence of congenital heart disease. J Epidemiol Community Health 1991;45: 299-301.
Zhang J, Ratcliffe JM. Paternal smoking and birthweight in Shanghai. Am J Public Health 1993;83:207-2IO.
Zhang X, Lv CC, Tian J, Miao RJ, Xi W, Hertz-Picciotto I, Qi L. Prenatal and perinatal risk factors for autism in China. J Autism Dev Disord 2010;40:I3|I-I321.
Zhu JL, Madsen KM, Vestergaard M, Basso O, Olsen J. Paternal age and preterm birth. Epidemiol 2005a; 16:259-262.

Zhu JL, Madsen KM, Vestergaard M, Olesen AV, Basso O, Olsen J. Paternal age and congenital malformations. Hum Reprod 2005b;20:3I73-3I77.
Zhu JL, Vestergaard M, Madsen KM, Olsen J. Paternal age and mortality in children. Eur J Epidemiol 2008;23:443-447.
Zhu JL, Olsen J, Liew Z, Li J, Niclasen J, Obel C. Parental smoking during pregnancy and ADHD in children: the Danish national birth cohort. Pediatrics 2014;134: e382-e388.
Zorlu P, Ergor GUL, Tezic T, Duru F, Ertem U. Evaluation of risk factors in children with acute lymphoblastic leukemia. Turkish J Cancer 2002;32:5-1I.


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[^1]:    AOR, adjusted odds ratio; AHR, adjusted hazard ratio; LBW, low birth weight ( $<2500 \mathrm{~g}$ ); NND, neonatal death; OR, odds ratio; PTB, preterm birth ( $<37$ weeks); SGA, small for gestational age; SSB, small size at birth; VPTB, very PTB (<32 weeks)

[^2]:    AC, achondroplasia; APR, adjusted prevalence ratio; CHD, congenital heart defects; OI, osteogenesis imperfecta; PDA, persistent ductus arteriosus; TD, thanatophoric dysplasia

