Second to fourth digit ratio (2D:4D) and prostate cancer risk in the Melbourne Collaborative Cohort Study

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BACKGROUND: The ratio of the lengths of the index (2D) and ring (4D) fingers as measured by the ratio 2D : 4D has been suggested as a proxy indicator of prenatal androgen activity, with low 2D : 4D reflecting higher in utero testosterone exposure (Manning et al, 1998; McIntyre, 2006; Hönekopp and Watson, 2010). There are several lines of evidence indicating that 2D : 4D is affected by prenatal androgens (Breidlove, 2010), and that digit ratios are longitudinally stable (McIntyre et al, 2005; Trivers et al, 2006). Hormone exposure in early life has been implicated in the aetiology of numerous cancers (Potischman et al, 2005). Prostate cancer is a hormonally driven and regulated disease, but studies have failed to detect associations between a single measure of hormone levels in adulthood and prostate cancer risk (Roddam et al, 2008).

Two recent studies have aimed to assess whether 2D : 4D is associated with prostate cancer (Jung et al, 2011; Rahman et al, 2011). Both of these studies concluded that low 2D : 4D, and thereby high prenatal testosterone, is a marker of increased risk of prostate cancer. We examine whether 2D : 4D is associated with prostate cancer risk in a large sample of men participating in the Melbourne Collaborative Cohort Study (MCCS).

METHODS: A total of 6258 men participating in the Melbourne Collaborative Cohort Study had 2D : 4D assessed. Of these men, we identified 686 incident prostate cancer cases. Hazard ratios (HRs) and confidence intervals (CIs) were estimated for a standard deviation increase in 2D : 4D.

RESULTS: No association was observed between 2D : 4D and prostate cancer risk overall (HRs 1.00; 95% CIs, 0.92–1.08 for right, 0.93–1.08 for left). We observed a weak inverse association between 2D : 4D and risk of prostate cancer for age < 60, however 95% CIs included unity for all observed ages.

CONCLUSION: Our results are not consistent with an association between 2D : 4D and overall prostate cancer risk, but we cannot exclude a weak inverse association between 2D : 4D and early onset prostate cancer risk.

MATERIALS AND METHODS

The MCCS is a prospective cohort study of 41,514 people (17,045 men) recruited between 1990 and 1994, 99.3% of whom were aged 40–69 years. Details of the MCCS have been published previously (Giles and English, 2002). At a recent face-to-face follow-up conducted during 2003–2009, 6287 men had their hands photographed. The length of the index and ring fingers were measured from photocopies of the surface of the hand using digital Vernier calipers. The length of the index finger was divided by the length of the ring finger to obtain 2D : 4D, and Dr-l was defined as the difference between right and left 2D : 4D. Measurement was undertaken by a team of trained research assistants at Cancer Council Victoria.

Follow-up commenced at baseline attendance and ended at 30 June 2009, the date the participant left Australia, diagnosis of an unknown primary tumour, or death, whichever occurred first. Of 6287 men who attended follow-up, we excluded 29 with a pre-baseline diagnosis of invasive prostate cancer or unknown primary tumour, leaving 6258 men available for analysis. During a median follow-up of 16 years, 686 incident prostate cancer cases were identified via linkage to the Victorian Cancer Registry.

Statistical analysis

Overall hazard ratios (HRs) and 95% confidence intervals (CIs) for a standard deviation (s.d.) increase in 2D : 4D measures were obtained from Weibull models with age as the time axis.
Age-varying HRs were estimated by flexible parametric survival models (Lambert and Royston, 2009), incorporating restricted cubic splines with six knots to model the baseline hazard, and restricted cubic splines with one knot to allow the HR to vary with age. Knots were evenly spaced across the distribution of uncensored log survival times. Separate estimates of HRs and CIs for age <60 and ≥60 years were obtained by fitting Weibull models to time-split data. The data duplication method was used to fit competing risks Weibull models for tumour aggressiveness (Lunn and McNeil, 1995). Aggressive tumours were those with Gleason score >7 or stage IV, or having prostate cancer as cause of death. Separate models were fit for right and left 2D:4D and Dr-l, and all models were adjusted for country of birth and baseline smoking status, as these factors are associated with 2D:4D (Manning et al, 2000; Manning and Fink, 2011). Statistical analyses were performed using Stata 11.1 for Linux (Stata Corporation, College Station, TX, USA).

RESULTS

Characteristics of the study population are shown in Table 1. Non-cases were younger than cases on average, with median age at baseline of 54 years compared with 59 years. Seventy-one per cent of participants were born in Australia, New Zealand, or in the United Kingdom.

Estimates from Weibull models are presented in Table 2. We found no overall association between either left or right 2D:4D and prostate cancer risk (HRs 1.00; 95% CIs, 0.92–1.08 for right, 0.93–1.08 for left). Estimated age-varying HRs and 95% CIs are plotted in Figure 1. There is some indication that higher 2D:4D is associated with lower early onset prostate cancer risk. For instance, the HR for an increase of one standard deviation in 2D:4D and prostate cancer risk at an age of 55 years is approximately 0.80 (95% CI, 0.65–1.10) for both hands. Similarly, risk of prostate cancer for men older than 80 years appears to be slightly reduced with higher 2D:4D; however, CIs include unity for all observed ages. Splitting follow-up at an age of 60 years and estimating separate HRs for the two age brackets also suggests that higher 2D:4D might be associated with decreased prostate cancer risk before 60 years of age. The HRs for an increase of one standard deviation in 2D:4D and risk of prostate cancer for age <60 years were 0.91 (95% CI, 0.73–1.13) and 0.88 (95% CI, 0.71–1.10) for left and right hands, respectively. Estimated HRs did not vary substantially by tumour aggressiveness (Table 2). No associations were observed for Dr-l (data not shown).

DISCUSSION

We found no statistically discernable association between 2D:4D and risk of prostate cancer either overall or by age; however, we cannot exclude that there might be a small inverse association between 2D:4D and risk of prostate cancer diagnosed before the age of 60 years.

Two recent studies have examined whether 2D:4D is associated with prostate cancer risk. The first was a clinical cohort study of 366 Korean men presenting with lower urinary tract symptoms (Jung et al, 2011). This study found that the odds of being diagnosed with prostate cancer were significantly higher for men with low 2D:4D compared with high 2D:4D (OR 3.22, 95% CI, 1.33–7.78). The second was a large case–control study, which reported an inverse association between self-assessed right 2D:4D and odds of prostate cancer (OR for index finger longer than ring finger vs index finger shorter than ring finger 0.67, 95% CI, 0.57–0.80) (Rahman et al, 2011). This study also reported a remarkably strong association between 2D:4D and prostate cancer diagnosed before the age of 60 years (OR 0.13, 95% CI, 0.09–0.21).

Although we found no evidence for an association of the magnitude previously reported (and indeed no statistically discernable associations at all), the possibility that there might be a weak inverse association between 2D:4D and early onset prostate cancer risk is broadly consistent with the estimate reported by Rahman et al (2011), as well as the current understanding of the association between hormone exposure in utero and 2D:4D. There are several differences between the studies that could account for the differing results, from study design (population based cohort vs clinical cohort vs case–control), to digit measurement method (photocopy vs direct vs self-reported assessment).

Our study has the advantage of being a large, population-based cohort with complete follow-up in terms of cancer diagnosis. Digit measurements were made with a high degree of reliability by trained research assistants. A disadvantage of our study is that not
Figure 1  Time‐varying HRs and 95% CIs from flexible parametric time to event models, adjusted for ethnicity and baseline smoking status.

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Author contributions

DCM planned and performed the statistical analysis and drafted the manuscript. JTM provided advice regarding the measurement of digit ratios and critically revised the manuscript. GGG, JLH, DRE and GS conceived the study, participated in the planning of the statistical analysis and critically revised the manuscript. All authors read and approved the final manuscript.

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Our analysis does not confirm the strong inverse association between 2D:4D and risk of prostate cancer previously reported. We cannot exclude the possibility that high 2D:4D is associated with lower risk of early onset prostate cancer. High 2D:4D is a marker of low in utero testosterone exposure, and thus hormone activity, early in development, might impact upon later risk of prostate cancer. Further research is required to clarify any association between 2D:4D and prostate cancer, especially for younger men.

all participants attended follow‐up, and thus we do not have 2D:4D available for every participant. If 2D:4D is associated with disease severity, and if disease severity is in turn associated with attendance, estimated associations could be biased. As there is evidence that 2D:4D is stable over time (McIntyre et al, 2006; Trivers et al., 2006), we consider it appropriate to analyse these data prospectively, despite the retrospective collection of 2D:4D. Another limitation of our study is the relatively small number of early onset prostate cancer cases, leading to limited power to detect associations between 2D:4D and prostate cancer risk for younger men.

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