Figure S1. Flowchart of exclusions of A: participants and B: metabolites.

The number of samples contributed to the data set twice because three controls became cases and five controls were controls for two cases each.

There were 2162 samples but 8 samples contributed to the data set twice because three controls became cases and five controls were controls for two cases each.
Figure S2. Partial correlation coefficients between metabolite and total PSA concentrations at baseline. Correlations between total PSA and metabolites were based on 764 controls for which total PSA was available, while correlations between metabolites included 1077 controls. The analyses were based on log-transformed data and were adjusted for age at blood collection (<55; 55-59; 60-64; 65-69; ≥70 years), body mass index (fourths; unknown) and study centre.
Figure S3. Statistical significance of the associations between metabolite concentrations and risk of high grade prostate cancer. High grade prostate cancer was tumours with Gleason score ≥8 or coded as undifferentiated tumours. The analysis included 124 matched case control sets. Statistical significance was plotted as −log_{10}(p-values). The dashed line represents conventionally statistical significance at α = 0.05; no associations were statistical significance after controlling the false discovery rate at α = 0.05 (Benjamini-Hochberg). Filled circles represent positive associations and unfilled circles represent inverse associations. The p-values were derived from a conditional logistic regression using log-metabolite concentration as a continuous variable and adjusting for exact age (continuously), body mass index (fourths; unknown), smoking (never; past; current; unknown), alcohol intake (<10; 10-19; 20-39; ≥40 g of alcohol per day; unknown), education (primary or none; secondary; degree level; unknown) and marital status (married or cohabiting; not married or cohabiting; unknown).
Figure S4. Statistical significance of associations between metabolite concentrations and risk of advanced stage prostate cancer. Advanced stage prostate cancer was defined as tumours with TNM score of T3-4 and/or N1-3 and/or M1, or coded as advanced. The analysis included 208 matched case control sets. Statistical significance was plotted as $-\log_{10}(p)$-values. The dashed and the dotted lines represent conventionally statistical significance and statistical significance after allowing for multiple testing using a false discovery rate controlling procedure (Benjamini-Hochberg), respectively, both at $\alpha = 0.05$. Filled circles represent positive associations and unfilled circles represent inverse associations. The p-values were derived from a conditional logistic regression using log-metabolite concentration as a continuous variable and adjusting for exact age (continuously), body mass index (fourths; unknown), smoking (never; past; current; unknown), alcohol intake (<10; 10-19; 20-39; ≥40 g of alcohol per day; unknown), education (primary or none; secondary; degree level; unknown) and marital status (married or cohabiting; not married or cohabiting; unknown).
Figure S5. Statistical significance of the associations between metabolite concentrations and risk of aggressive prostate cancer. Aggressive prostate cancer was tumours with TNM score of T4 and/or N1,3 and/or M1. The analysis included 115 matched case control sets. Statistical significance was plotted as $-\log_{10}(p)$-values. The dashed line represents conventionally statistical significance at $\alpha = 0.05$; no associations were statistical significance after controlling the false discovery rate at $\alpha = 0.05$ (Benjamini-Hochberg). Filled circles represent positive associations and unfilled circles represent inverse associations. The $p$-values were derived from a conditional logistic regression using log-metabolite concentration as a continuous variable and adjusting for exact age (continuously), body mass index (fourths; unknown), smoking (never; past; current; unknown), alcohol intake (<10; 10-19; 20-39; ≥40 g of alcohol per day; unknown), education (primary or none; secondary; degree level; unknown) and marital status (married or cohabiting; not married or cohabiting; unknown).
Figure S6. Statistical significance of associations between metabolite concentrations and risk of death from prostate cancer. The analysis included 127 matched case control sets. Statistical significance was plotted as $-\log_{10}(p$-values). The dashed line represents conventionally statistical significance at $\alpha = 0.05$; no associations were statistical significance after controlling the false discovery rate at $\alpha = 0.05$ (Benjamini-Hochberg). Filled circles represent positive associations and unfilled circles represent inverse associations. The p-values were derived from a conditional logistic regression using log-metabolite concentration as a continuous variable and adjusting for exact age (continuously), body mass index (fourths; unknown), smoking (never; past; current; unknown), alcohol intake ($<10$; $10-19$; $20-39$; $\geq 40$ g of alcohol per day; unknown), education (primary or none; secondary; degree level; unknown) and marital status (married or cohabiting; not married or cohabiting; unknown).