symptoms or visible discharge and ≥5 PMNs/high powered field (HPF). Absence of CT, MG, adenoavirus, and HSV was considered as idiopathic NGU. Men without NGU had no urethral symptoms, no discharge, and <5 PMNs/HPF. Broad-range 16S rRNA gene PCR with deep sequencing was used to characterize the urethral microbiota. Compositional lasso analysis of bacterial taxa was conducted to identify associations between bacteria and NGU; beta coefficients (β) giving change in probability of NGU per log2 change in relative abundance are reported.

Results Of 434 (199 MSM, 235 MSW) urine samples, 330 yielded sequence data. NGU+ men were less likely to yield sequence data (70% vs 84%, Fisher’s p = 0.001). Of 328 men with ≥1000 sequence reads/sample, 95 MSM (44 NGU+) and 143 MSW (46 NGU+) were negative for CT, MG, adenoavirus, and HSV. Higher relative abundances of Haemophilus influenzae (β = 0.0139) and Mycoplasma penetrans (β = 0.0095) were positively associated with idiopathic NGU in MSM, while H. influenzae was positively associated with idiopathic NGU in MSW (β = 0.0184). The model also identified bacterial species that were negatively associated with NGU in MSM and MSW. Notably, Lactobacillus iners was negatively associated with idiopathic NGU in MSW (β = −0.0006) but not MSM.

Conclusion Different bacterial species are associated with NGU in MSM and MSW. We identified two bacterial species infrequently detected in the male urethra as positively associated with NGU.

Disclosure No significant relationships.

P524 GAYS, GOVERNMENT AND BIG DATA: SHOULD ROUTINE HEALTH RECORDS INCLUDE SEXUAL ORIENTATION?

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Background Sexual orientation minorities continue to experience poorer outcomes in sexual health, mental health and addictions. Despite clear information needs, routine data identifying gay, lesbian and bisexual (GLB) individuals are seldom collected by governments, rendering these populations invisible. In New Zealand (NZ), everyone is assigned a unique National Health Index (NHI) number used across all health systems to improve clinical and public health decision-making. In 2017 the NZ Ministry of Health proposed adding sexual orientation and other fields to NHI, however sexual orientation and other fields to NHI, however sexual orientation was rejected after consultation. We sought to better understand these viewpoints.

Methods We used the Official Information Act (OIA) to request the complete list of submitters, their support or opposition for adding sexual orientation and other fields to NHI, however sexual orientation was rejected after consultation. We sought to better understand these viewpoints.

Results 130 submissions were received. Overall 27 supported sexual orientation in NHI and 35 were opposed, the remainder being neutral or conditional. Support by grouping (high to low) was: Academic (100%); Government (56%); Unspecified (44%); NGO (40%); Health (36%) and Data firm (25%). Supportive reasons included: service planning; evidence-based policy; equity; GLB-specific health delivery (e.g. HPV vaccines, HIV and STI screening, pre-exposure prophylaxis); normalisation; health workforce development. Opposing reasons included: sexual orientation being irrelevant to clinical decision-making; classification challenges (e.g. sexual orientation fluidity); data quality; privacy; discrimination.

Conclusion Barriers to sexual orientation data collection include practical concerns but also well-meaning paternalism and heteronormative assumptions. Better information about NHS uses and protections, data governance, and healthcare service obligations to GLB communities, may improve support.

Disclosure No significant relationships.

P525 PREVALENCE OF MYCOPLASMA GENITALIUM BY ANATOMICAL SITE IN MEN WHO HAVE SEX WITH MEN: A SYSTEMATIC REVIEW AND META-ANALYSIS

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Background With the current debate over testing and screening for Mycoplasma genitalium (MG) in various populations, more information on the prevalence of MG is needed particularly in populations at high risk of sexually transmitted infections, such as men who have sex with men (MSM). We assessed the available data on the prevalence of MG in MSM across three anatomical sites: the urethra, pharynx and rectum.

Methods Ovid Medline, PubMed, Embase were searched for all peer-reviewed studies published until 1st June 2018 (in addition to conference proceedings from 2015), that reported prevalence of MG (using nucleic acid amplification testing) in the urethra, pharynx and/or pharynx in at least 50 MSM. Data were extracted by anatomical site, symptom and HIV status. Pooled estimates (95% confidence intervals [CIs]) were calculated using random effects meta-analysis. Subgroup analyses were performed to assess heterogeneity between studies.

Results Forty-six studies met inclusion criteria. The overall prevalence of MG at any site was 5.8% (4.5–7.3%, I2 = 95.0%). MG prevalence was 4.6% (3.0–6.4%, I2 = 94.4%) in the urethra, 6.1% (4.5–7.9%, I2 = 89.0%) in the rectum, and 1.0% (0.0–5.1%, I2 = 96.0%) in the pharynx. Poole estimates of MG prevalence were higher among HIV-positive compared with HIV-negative men (9.0% [5.2–13.4%, I2 = 90.7%] versus 5.7% [3.5–8.2%, I2 = 93.1%, p = 0.019], and among asymptomatic men compared to symptomatic men (9.2% [6.2–12.7%, I2 = 87.3%] versus 4.0% [2.3–6.2%, I2 = 90.7], p = 0.003).

Conclusion MG is commonly detected in MSM, particularly in the urethra and rectum. Prevalence was highest in HIV