with a rapid test and a second test using an enzyme immunoassay (EIA) was developed (average turnaround time of 2 weeks). Given the changes in the market since 2003, we present results of a second rapid test validation to update the Guatemalan algorithm.

**Methods** Validation was performed in 2012–2013, evaluating a total of 10 rapid tests in two phases: 1) with serum, HIV-negative samples were obtained from blood banks, and positive samples from HIV care units; 2) with whole blood, negative samples were obtained from antenatal care services and positive samples from HIV care units. We tested 452 serum samples [244 HIV+, 208 HIV-] and 432 whole blood samples (10 HIV positive, 422 HIV negative) using two EIA (Roche ELECSYS HIV Combi and Abbott AXSYM HIV Ag/Ab Combo). Discordant results were evaluated with HIV-1 Western Blot. For fourth generation rapid tests, only antibody was evaluated. Data was analysed using OpenEpi.

**Results** Six HIV rapid tests were included for both phases. All tests (except Double Check Gold) reported sensitivity higher than 99% and specificity higher than 95%. Determine HIV 1/2 and HIV Ag/Ab reported the highest results for sensitivity (100%). Hexagon HIV (100%) and Accu-Tell (99.5%) reported the highest results for specificity. All rapid tests that also included HIV-2 detection, showed cross-reactivity, ranging from 37% of HIV cases for Anarapid HIV 1/2/O Tri-line to 15% for Rapid HIV 1/2/O Tri-Line.

**Conclusion** Our results show that Determine HIV½ and HIV Ag/Ab should be recommended as a screening test while Hexagon HIV and Accu-Tell should be recommended as second test. Rapid test validation provides valuable information for on-site confirmation of reactive results improving diagnosis turnaround time.

**Disclosure of interest statement** We declare that we have no conflicts of interest.

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**P17 - HIV testing, treatment and care**

**P17.01 PERFORMANCE EVALUATION OF THE GENEXPERT HIV-1 QUANT ASSAY FOR DETECTION OF HIV-1 IN PLASMA**

L McNally, A Carrera, J Sherring, PH Cunningham. NSW State Reference Laboratory for HIV, St Vincent’s Hospital Sydney Limited, Kirby Institute UNSW

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**Introduction** The HIV-1 Quant Assay performed on the GeneXpert® Instrument Systems, is designed for the rapid quantitation of HIV-1 in human plasma with an analytical range of 40 to 10,000,000 copies/mL for HIV-1 Group M subtypes A, C, D, AE, F, G, H, AB, AG, J, K and Groups N and O. Testing is performed in a single-use disposable GeneXpert cartridges that hold the real-time reverse transcriptase polymerase chain reaction (RT-PCR) reagents and host the RT-PCR processes. This study assessed the performance of the system in routine plasma.

**Methods** To date, a total of 130 plasma samples have been tested over the analytical range and compared to a benchmark real time PCR system. Seventy four samples (56.9%) were of a known subtype comprising of subtype B (37.6%), AE (7.7%), C (4.6%) AG (1.5%) and mixed (4.6%). Additional samples consisting of an external quality control samples run over multiple days, and samples with HIV-1 RNA not detected or below the lower limit of were also tested to assess performance.

**Results** Overall the HIV-1 Quant Assay performed on the GeneXpert® Instrument Systems correlated with the routine analytical platform (r² = 0.9333). Samples ranged undetectable (16, 8.8%), below the benchmark test lower limit of detection (<20 cpy/ml) (10, 7.9%), low range (20–5,000) (43, 33.1%), medium range (5,000–50,000) (24, 18.5%) and high range (>50,000 cpy/ml) (29, 22.3%). Thirteen samples (10%) were invalid as a result of insufficient sample. Samples in the lower analytical range <1,000 cpy/ml showed little variance when compared with the Roche (CÂP/CTM) assay using Bland-Altman correlation analysis. Reproducibility was assessed in the high, medium and low range within 1–2SD of mean. Sixteen replicates of a commercial external control showed very good reproducibility.

**Conclusion** The HIV-1 Quant Assay performed on the GeneXpert® Instrument Systems correlated with a commonly used HIV RNA test in plasma and offered significant workflow advantages. The system has a small footprint and requires no further consumables other that the single-use test cartridges. Further studies are planned to fully assess the assay performance.

**Disclosure of interest statement** No conflict of interest to declare.

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**P17.02 ORAQUICK® IN-HOME HIV TEST KIT IN PERU: AVAILABILITY AND ACCEPTABILITY AMONG MEN WHO HAVE SEX WITH MEN AND TRANSGENDER WOMEN**

1Maria Jose Bustamante*, 1Kelka A Konda, 2Segundo R León, 3Gino Calvo, 4Javier Salvaterra, 5Brandon Brown, 6Carlos F Caceres, 7Jeffrey D Klausner. 8Unit of Health, Sexuality and Human Development, and Laboratory of Sexual Health, Universidad Peruana Cayetano Heredia, Lima, Peru; 9Program in Global Health, Department of Medicine, University of California Los Angeles, Los Angeles CA, USA; 10Epicentro Salud, Lima, Peru; 11Barton Health Center, Health Directorate of Callao, Lima, Peru; 12Program in Public Health, University of California Irvine, Irvine CA, USA

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**Background** In Peru an estimated 70% of people who are HIV positive do not know their status. Knowing one’s HIV status is critical in HIV prevention. Oraquick® in-home HIV test is the first rapid HIV self-test U.S FDA-approved for home use. We aimed to assess its availability and the willingness of men who have sex with men (MSM) and transgender women (TW) to use it in Peru.

**Methods** Four Pharmacy chains in Peru were surveyed to ascertain commercial availability of the Oraquick® in-home HIV test kits. High-risk MSM and TW who attended either of two STI clinics in Lima from June 2013 to May 2014 were surveyed. Data on demographics and willingness to use Oraquick® IN HOME HIV TEST kit were collected using an interviewer administered computer-based questionnaire.

**Results** The Oraquick® Rapid HIV-1/2 test kit was available for purchase for home use by 4 (100%) pharmacy chains, 3 in Lima and one in northern Peru. The average test kit cost was 54 soles (18 USD); kits were available to clients 18 years or older for over-the-counter purchase. Of the 137 interviewed survey participants, 85% (n = 117) reported they would use a rapid home HIV self-test at least twice yearly. Respondents reported willingness to pay up to 21 soles (approximately 7 USD) for the test and in the event of positive results, to do the confirmatory blood test in a clinic. Also, 78% (n = 117) of participants reported being comfortable getting an HIV self-test kit by mail or for home use from a clinic.

**Conclusion** Our findings show the potential utility for home HIV self-testing to enhance HIV serostatus awareness in Peru.
Further studies are needed on utilisation and linkage to care and prevention services.

**P17.03** CONVENIENT HIV TESTING SERVICE MODELS COMBINED WITH RAPID TESTING ARE ATTRACTING PREVIOUSLY UNTESTED GAY AND BISEXUAL MEN

V Knight*, H Wand, J Gray, P Keen, A McNulty, G Guy, Sydney Sexual Health Centre, South East Sydney Local Health District, Sydney, 2000, Australia; The Kirby Institute, University of New South Wales, Sydney, NSW, 2052, Australia; ACON Health, 414 Elizabeth Street Sunny Hills, Sydney, NSW, 2010, Australia; School of Public Health and Community Medicine, University of NSW, Kensington, NSW, 2052, Australia

**Introduction** HIV testing is a cornerstone of the treatment as prevention (TasP) approach. In Australia, gay and bisexual men (GBM) account for the majority of HIV diagnoses each year, but less than a quarter are testing at the recommended frequency. In response, a range of new service delivery models were introduced in Sydney over the past few years. We assessed which HIV testing service delivery models were more likely to attract GBM who had never previously tested or were testing infrequently.

**Methods** We compared demographics, risk behaviour and HIV testing history among new GBM clients attending three different HIV testing service models (fast-track Xpress clinic, fixed-site community-based service and time-limited community-based shopfront) between August 2013 and May 2014. All services offered HIV rapid testing. We used multivariate regression to assess factors (including service model) associated with being untested or infrequent testers (not tested within the past 12 months).

**Results** Overall, 1704 new GBM attended the services; 19% were untested and 41% were infrequent testers. Across the services, there were significant differences in demographics, risk behaviour and past HIV testing history. The overall HIV seropositivity was 1.2% (95% CI: 0.8%–1.9%) and STI positivity was 12.4% (95% CI: 11.6–17.2) with no significant differences across services. Factors independently associated with being untested were attendance at the two community sites, younger age, being born in Asia, living in North Sydney, being bisexual and reporting fewer male sexual partners. Factors independently associated with infrequent testers were attending the fast track Xpress clinic, being older, being born in Asia and reporting fewer male partners.

**Conclusion** The findings show the two community sites reached more untested men and the fast-track clinic model more infrequent testers but the HIV/STI diagnosis rate was consistent across services, indicating that all three testing models are important to increase HIV testing among GBM.

**Disclosure of interest statement** Vickie Knight is supported by an Australian Postgraduate Association scholarship. No pharmaceutical grants were received in the development of this study.

**P17.04** THE RELATIONSHIP BETWEEN HIV TESTING FREQUENCY AND HIV RISK PERCEPTION AMONG PERUVIAN MSM AND TRANSGENDER WOMEN

Sk Vargas*, KA Konda, LH Leon, GC Calvo, HI Salvatierra, B Brown, DJ Klauser, CF Caceres. Unit of Health, Sexuality and Human Development, and Laboratory of Sexual Health, Universidad Peruana Cayetano Heredia, Lima, Peru; University of California, Los Angeles, CA, USA; Epicentro, Lima, Peru; Alberto Barton Health Center, Health Directorate of Callao, Lima, Peru; University of California, Irvine, CA, USA

**Introduction** HIV in Peru is concentrated among men who have sex with men (MSM) and transgender women (TW). Over 70% of HIV+ MSM/TW ignore their status, as HIV testing frequency is low, delaying diagnosis and treatment. We aimed to assess the relationship between HIV testing frequency and perceived HIV risk among MSM/TW.

**Methods** This analysis included baseline data from MSM/TW who reported a negative or unknown HIV status from a STI clinic cohort of MSM/TW at high risk in Lima. The behavioural survey assessed HIV risk perception (high, moderate, low) and prior testing frequency (testing was defined as frequent if it had occurred at least biannually). For HIV diagnosis, we used a 3rd generation rapid test, a 4th generation ELA, and Western Blot confirmation. Chi-square tests and multivariable regression were used to estimate adjusted prevalence ratios (aPRs).

**Results** Among eligible subjects (243 MSM and 67 TW), 122 (39%) reported frequent prior HIV testing, while HIV risk was perceived as high, moderate and low by 72 (23%), 148 (48%) and, 90 (29%) participants, respectively. Frequent HIV testing was more prevalent among participants reporting low/no HIV risk (aPR: 1.62; CI: 1.20–2.0) and among those reporting previous syphilis infection (aPR: 1.58; CI: 1.22–2.04); but was less prevalent among participants reporting recent condomless anal intercourse (aPR: 0.67; CI: 0.51–0.90). Baseline HIV status and HIV risk perception were not associated: HIV prevalence was 11%, 13% and 10% among those reporting high, moderate and low/no HIV risk (chi-square p-value >0.1).

**Conclusion** Frequent HIV testing was more prevalent among MSM/TW reporting low/no HIV risk. Risk perception did not correlate with HIV status among previously undiagnosed subjects. WHO-recommended biannual HIV testing should be promoted, and factors potentially affecting risk perception and testing frequency (e.g. access, stigma) should be identified and addressed by HIV programming for MSM/TW in Peru.

**P17.05** HIV TEST REFUSALS AMONG BLACK AFRICANS ATTENDING SEXUALLY TRANSMITTED INFECTION CLINICS IN ENGLAND, 2013

H Mohammed*, M Furegato, A Nardone, G Hughes. HIV & STI Department, Public Health UK

**Introduction** Black Africans comprise the second largest group affected by HIV in England, but it is estimated that 34% are