Example: chlamydia in England, 2012

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1 Example: Chlamydia in England, 2012

This example illustrates a method for using chlamydia surveillance data to estimate prevalence. Surveillance data on chlamydia testing and diagnosis rates in England in 2012 were downloaded from: http://www.chlamydiascreening.nhs.uk/ps/data.asp (downloaded 9 February 2016).

		Men	Wome	en		
	15-19 years	20-24 years	Total	15-19 years	20-24 years	Total
Population	1685620	1833395	3519015	1600686	1788156	3388842
Tests	232668	334240	566908	520358	685538	1205896
Diagnoses	15213	33174	48387	42874	45227	88101

Data on sexual behaviour from the third National Study of Sexual Attitudes and Lifestyles (Natsal-3) are available from the UK data service: https://www.ukdataservice.ac.uk/ (downloaded 23 September 2015). These were used to infer 95% confidence intervals for the proportions of men and women, aged 16-19 and 20-24, who were sexually active (see the accompanying R script; note that no 15-year-olds were recruited to Natsal-3). These 95% confidence intervals were in turn used to derive beta-distribution priors for the proportion sexually active within each sex and age group.

1.1 Sampling for testing and diagnosis rates

```
In [1]: import numpy as np
        from numpy import *
        from scipy.stats import beta
        from scipy.optimize import fsolve
        ########################
        # parameters of beta distributions representing the proportion of the population sexually
        # active, by sex and age group
        ############################
        # men. 16-19
        [alpha_m_16_19, beta_m_16_19] = fsolve(
            lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
            - (0.6747424, 0.741327698),
            [1,1]
        # men, 20-24
        [alpha_m_20_24, beta_m_20_24] = fsolve(
            lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
            (0.8844970, 0.933759842).
            [1,1]
            )
        # men, 16-24
        [alpha_m_16_24, beta_m_16_24] = fsolve(
            lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
```

```
- (0.8023836019, 0.843403825),
   [1,1]
   )
# women, 16-19
[alpha_f_16_19, beta_f_16_19] = fsolve(
   lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
    - (0.6583593, 0.723554878),
   [1,1]
   )
# women, 20-24
[alpha_f_20_24, beta_f_20_24] = fsolve(
   lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
   - (0.8904135, 0.934417684),
   [1,1]
   )
# women, 16-24
[alpha_f_16_24, beta_f_16_24] = fsolve(
   lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
   - (0.7998634469, 0.837979601),
   [1,1]
```

Next, sample from distributions for the probability of being sexually active, the size of the sexually active population and the testing and diagnosis rates per person per year.

```
In [2]: from scipy.stats import gamma
        from numpy.random import normal
        rs = random.RandomState(12345)
        n_sample = 10000
        # sexually-active populations:
        p_active_m_16_19 = rs.beta(alpha_m_16_19, beta_m_16_19, size=n_sample) # 16-19 yo only
        pop_active_m_15_19 = rs.binomial(1685620, p_active_m_16_19, size=n_sample)
        p_active_m_20_24 = rs.beta(alpha_m_20_24, beta_m_20_24, size=n_sample) # 20-24 yo only
        pop_active_m_20_24 = rs.binomial(1833395, p_active_m_20_24, size=n_sample)
        p_active_m_16_24 = rs.beta(alpha_m_16_24, beta_m_16_24, size=n_sample) # 16-24 yo only
        pop_active_m_15_24 = rs.binomial(3519015, p_active_m_16_24, size=n_sample)
        p_active_f_16_19 = rs.beta(alpha_f_16_19, beta_f_16_19, size=n_sample) # 16-19 yo only
        pop_active_f_15_19 = rs.binomial(1600686, p_active_f_16_19, size=n_sample)
        p_active_f_20_24 = rs.beta(alpha_f_20_24, beta_f_20_24, size=n_sample) # 20-24 yo only
        pop_active_f_20_24 = rs.binomial(1788156, p_active_f_20_24, size=n_sample)
        p_active_f_16_24 = rs.beta(alpha_f_16_24, beta_f_16_24, size=n_sample) # 16-24 yo only
        pop_active_f_15_24 = rs.binomial(3388842, p_active_f_16_24, size=n_sample)
        # testing and diagnosis rates, per person per year
        \texttt{test\_rate\_m\_15\_19} = \texttt{rs.gamma}(232668, 1, \texttt{size=n\_sample})/\texttt{pop\_active\_m\_15\_19}
        test_rate_m_20_24 = rs.gamma(334240, 1, size=n_sample)/pop_active_m_20_24
        test_rate_m_15_24 = rs.gamma(566908, 1, size=n_sample)/pop_active_m_15_24
        diag_rate_m_15_19 = rs.gamma(15213, 1, size=n_sample)/pop_active_m_15_19
        diag_rate_m_20_24 = rs.gamma(33174, 1, size=n_sample)/pop_active_m_20_24
        diag_rate_m_15_24 = rs.gamma(48387, 1, size=n_sample)/pop_active_m_15_24
        diag_rate_f_15_19 = rs.gamma(42874, 1, size=n_sample)/pop_active_f_15_19
        diag_rate_f_20_24 = rs.gamma(45227, 1, size=n_sample)/pop_active_f_20_24
        diag_rate_f_15_24 = rs.gamma(88101, 1, size=n_sample)/pop_active_f_15_24
        test_rate_f_15_19 = rs.gamma(520358, 1, size=n_sample)/pop_active_f_15_19
        test_rate_f_20_24 = rs.gamma(685538, 1, size=n_sample)/pop_active_f_20_24
        test_rate_f_15_24 = rs.gamma(1205896, 1, size=n_sample)/pop_active_f_15_24
In [3]: print percentile(test_rate_m_15_24,50)
        print percentile(diag_rate_m_15_24,50)
```

1.2 Sampling natural history, behavioural and other parameters

Priors for the proportion of incident infections which are asymptomatic and the test performance were taken directly from published studies. The proportion of incident infections which are asymptomatic is not known, so we use estimates of the proportion of *prevalent* infections asymptomatic. Note that these provide an upper bound because treatment seeking by symptomatic cases will deplete the symptomatic pool.

```
In [4]: # test performance
```

```
# Horner J. Clin. Microbiol (2005): 32 of 32 infected samples tested +ve
p_true_pos_m = rs.beta(32+1, 0+1, size=n_sample)
# Horner J. Clin. Microbiol (2005): 2 of 952 uninfected samples tested +ve
p_false_pos_m = rs.beta(2+1, 950+1, size=n_sample)
# Low Health Technol Assess (2007): 129 of 141 infected samples tested +ve
p_true_pos_f = rs.beta(129+1, 12+1, size=n_sample)
# Low Health Technol Assess (2007): 4 of 2327 uninfected samples tested +ve
p_false_pos_f = rs.beta(4+1, 2323+1, size=n_sample)
```

1.2.1 Rate of treatment seeking by symptomatic cases

We use a Metropolis-Hastings algorithm to sample for the rate of treatment following onset of symptoms, assuming a constant hazard of treatment beginning with the onset of symptoms. Data consist of the estimated proportion of GUM clinic patients with symptoms whose symptoms had started less than one, 1-2, 2-4, 4-6 and more than 6 weeks previously (Mercer et al., Sex. Transm. Infect. 83:400-405; 2007).

Proportion					
	Estimate	95% Confidence Interval			
< 1 week	26.7%	(14.4, 44.2)%			
7-13 days	14.4%	(6.1, 30.2)%			
14-27 days	20.8%	(13.3, 31.0)%			
4-6 weeks	16.6%	(8.5, 29.9)%			
>6 weeks	21.5%	(5.5, 56.4)%			

```
In [5]: # function for calculating likelihood of multinomial data
       %run multinomial_pmf.py
In [6]: # Find beta distributions corresponding to 95% CIs reported in
        # Mercer Sex. Transm. Infect. (2007) (see table above).
        a = empty(5)
       b = empty(5)
        # < 1 week
        [a[0], b[0]] = fsolve(
            lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
            - (0.144, 0.442),
            [1,1]
        # 7-13 days
        [a[1], b[1]] = fsolve(
            lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
            - (0.061, 0.302),
            [1,1]
            )
        # 14-27 days
        [a[2], b[2]] = fsolve(
            lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
```

```
- (0.133, 0.310),
            [1,1]
            )
        # 28-41 days
        [a[3], b[3]] = fsolve(
            lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
            - (0.085, 0.299),
            [1,1]
            )
        # 42 days and over
        [a[4], b[4]] = fsolve(
            lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
            - (0.055, 0.564).
            [1,1]
In [7]: # Metropolis-Hastings to get a sample for rate of treatment
        att_symp = empty(n_sample+1000) # testing rate per person per year. Allow 1000 extra samples for burn-in
        11 = empty(n_sample+1000) # log-likelihood
       props = empty([n_sample+1000, 5]) # simulated data, for posterior predictive check
       old = 0.04 # starting sample value
       new = 0.04 # starting sample value
        # simulate probabilities corresponding to data
        # proportion expected in each time window
       tps = array([0., 7., 14., 28., 42., Inf])
        simp_old = exp(-old*tps[:5]) - exp(-old*tps[1:])
       simp_new = exp(-new*tps[:5]) - exp(-new*tps[1:])
       while i < n_sample+1000: # to do samples for p_test_symp
            new = rs.normal(old, 0.05) # generate a sample from normal distribution
            if new < 0:
                att_symp[i] = old # reject
                11[i] = -1e10
            else:
                simp_old = exp(-old*tps[:5]) - exp(-old*tps[1:])
                simp_new = exp(-new*tps[:5]) - exp(-new*tps[1:])
                if sum(simp_new > 0) != len(tps) - 1:
                    att_symp[i] = old # reject
                    11[i] = -1e10
                ٠ مه [م
                    # simulate probabilities corresponding to the data
                    log_ratio = \
                        sum(beta.logpdf(simp_new, a, b, loc=0, scale=1)) \
                        - sum(beta.logpdf(simp_old, a, b, loc=0, scale=1))
                    if log(rs.uniform(0,1)) < log_ratio:</pre>
                        att_symp[i] = new # accept
                        11[i] = sum(beta.logpdf(simp_new, a, b, loc=0, scale=1))
                        old = new
                        acc = acc+1
                    else:
                        att_symp[i] = old # reject
                        11[i] = sum(beta.logpdf(simp_old, a, b, loc=0, scale=1))
            props[i] = simp_old
        att_symp = att_symp[1000:] # remove burn-in samples
        11 = 11[1000:] # log-likelihood
```

```
print acc/(n_sample+1000) # print the proportion of samples accepted
       print mean(att_symp)*365.25
       print array(percentile(att_symp, [2.5, 97.5]))*365.25
        att_symp = att_symp*365.25 # convert rate from day^-1 to year^-1
0.226545454545
14.4054933827
[ 8.59839927 22.22498957]
In [8]: # diagnostics and posterior predictive checks
        import matplotlib.pyplot as plt
        %matplotlib inline
       from numpy.random import multinomial
       fig = plt.figure(figsize = (10,10))
        ax1 = fig.add_subplot(221)
       ax1.plot(att_symp, alpha=0.5)
        ax2 = fig.add_subplot(222)
        ax2.plot(range(43), median(att_symp/365.25)*exp(-median(att_symp/365.25)*array(range(43))),'b')
        \#plt.plot(range(50),percentile(att\_symp,~2.5)*exp(-percentile(att\_symp,~2.5)*array(range(50))), 'b--')
        #plt.plot(range(50), percentile(att_symp, 97.5)*exp(-percentile(att_symp, 97.5)*array(range(50))), 'b--')
        #ax2.set_ylim([0,0.1])
        ax2.set_xlim([0,50])
        ax2.errorbar([3.5,10.5,21,35, 46],
                    [0.267/7, 0.144/7, 0.208/14, 0.166/14, 0.215/10],
                    abs(array([[0.144/7, 0.061/7, 0.133/14, 0.085/14, 0.055/10],
                               [0.442/7, 0.302/7, 0.310/14, 0.299/14, 0.564/10]]
                              ) - array([0.267/7, 0.144/7, 0.208/14, 0.166/14, 0.215/10])
            ), color = 'r', fmt='.')
        ax2.plot([0,7,7,14,14,28,28,42], repeat(percentile(props[:,:4],50,0)/array([7,7,14,14]),2), 'b--')
        ax2.plot([42,50], repeat(percentile(props[:,4],50,0)/array([10]),2), 'b--')
        ax2.fill_between(
            [0,7,7,14,14,28,28,42,42,50],
            repeat(percentile(props, 2.5, 0) / array([7,7,14,14,10]), 2),
            repeat(percentile(props, 97.5,0)/array([7,7,14,14,10]),2),
            alpha=0.5
        )
        ax1.set_xlabel('Sample')
        ax1.set_ylabel('Rate of seeking treatment (year^-1)')
        ax2.set_xlabel('Days since onset of symptoms')
        ax2.set_ylabel('Proportion of patients surveyed')
Out[8]: <matplotlib.text.Text at 0x10ccae690>
```

The MCMC chain is illustrated in the left-hand panel, and seems to have converged well.

The right-hand panel shows the probability density of the time between onset of symptoms and attending the GUM clinic where patients were surveyed, to 42 days (solid blue line). The blue shaded area and dashed line show the central 95% and median of simulated histograms for waiting times to clinic, with bins corresponding to time windows reported in the data. The last bin contains all times longer than six weeks and has been divided by 10 (as opposed to the width of the window) to make it readable. For comparison, red error bars show the reported proportions of patients with treatment-seeking times within each time window (estimate and 95% CI), normalised to be on the same scale as the predictions (blue). The good predictive properties of the model are indicated by the agreement between the data, in red, and the posterior predictions in blue.

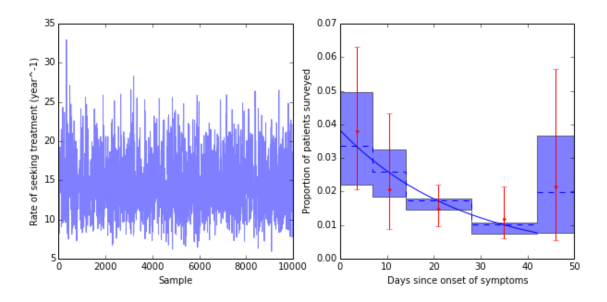


Figure 1: Diagnostic plots for MCMC sampling of treatment seeking rate in symptomatic patients. Left: MCMC chain. Right: posterior predictive check (for description, see text.)

1.2.2 Rate of spontaneous clearance of infection

Rates of spontaneous clearance of infection in men and women were sampled using MCMC and the STAN software (see accompanying R scripts, STAN model files and references), following the model presented by Price *et al.* in *Stat. Med.* **32**:1547-1560.

```
In [16]: import csv
        sc_m = empty(n_sample) # clearance rate per person per year
        with open('stan/chlamydia_two_exponentials_men.csv', 'rU') as m:
            reader = csv.reader(m)
            next(reader) # skip the header row
            for row in reader:
                sc_m[i] = row[0]
                i = i+1
        sc_f = empty(n_sample) # clearance rate per person per year
        with open('stan/chlamydia_two_exponentials_women.csv', 'rU') as f:
            reader = csv.reader(f)
            i=0
            next(reader) # skip the header row
            for row in reader:
                sc_f[i] = row[0]
                i = i+1
        h=plt.hist(sc_f, bins=50, histtype='step', normed=True, color='r')
        h=plt.hist(sc_m, bins=50, histtype='step', normed=True, color='b')
        plt.xlabel('Rate of spontaneous clearance (year$^{-1}$)')
        print 'Median (central 95% credible interval) for spontaneous clearance rate in men: \n \t', \
            percentile(sc_m, 50), percentile(sc_m, (2.5,97.5))
        print 'Median (central 95% credible interval) for spontaneous clearance rate in women:\n \t', \
            percentile(sc_f, 50), percentile(sc_f, (2.5,97.5))
Median (central 95% credible interval) for spontaneous clearance rate in men:
          0.395824502635 [0.058796284132470837, 1.2738753854360512]
Median (central 95% credible interval) for spontaneous clearance rate in women:
          0.725636286054 [0.59114145020803666, 0.87428940522833309]
```

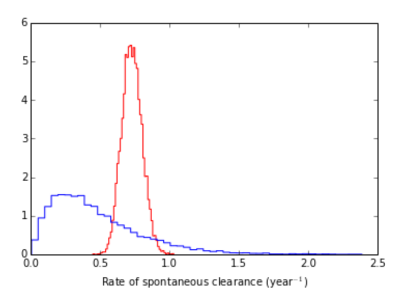


Figure 2: Sampled rates of spontaneous chlamydia clearance in men (blue) and women (red).

Finally, we infer the proportion of infections which are asymptomatic by calibrating to the Natsal-3 prevalence estimates in 16-25-year-old men and women.

```
In [10]: from scipy.stats import beta
         [alpha_prev_m, beta_prev_m] = fsolve(
             lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
             - (0.015, 0.034), # Natsal-3 prevalence in men
             [1,1]
             )
         prev_m = rs.beta(alpha_prev_m, beta_prev_m, size=n_sample)
         # generate samples for prevalence
         [alpha_prev_f, beta_prev_f] = fsolve(
             lambda x: array(beta.interval(0.95, x[0], x[1], loc=0, scale=1))
             - (0.022, 0.043), # Natsal-3 prevalence in women
             [1,1]
         prev_f = rs.beta(alpha_prev_f, beta_prev_f, size=n_sample)
In [11]: # This script also contains the functions linking observed tests, symptomatic/asymptomatic/toal diagnoses,
         # incidence, prevalence, screening and other model parameters
         # Running it takes a little while because of all the symbolic algebra
         %run test_diag_fun.py
In [12]: # incidence, screening and proportion of incident infections asymptomatic in men
         inc_m = np.zeros(n_sample)
         scr_m = np.zeros(n_sample)
         p_asymp_m = np.zeros(n_sample)
         for i in xrange(n_sample):
             def tmpfun(inc, scr, p_asymp):
                 [tr, dr] = test_diag_fun(
                     array([
                         inc,
                         1-p_asymp, # proportion of incident infections which are symptomatic
                         sc_m[i], # rate of self-clear
```

```
att_symp[i],
                        p_true_pos_m[i],
                       p_false_pos_m[i]
                       ]))
                prev = dyn_fun(
                    inc*p_asymp,
                    sc_m[i] + scr*p_true_pos_m[i],
                    inc*(1-p_asymp),
                    scr*p_true_pos_m[i] + att_symp[i]*p_true_pos_m[i]
                prev - prev_m[i])
            [inc_m[i], scr_m[i], p_asymp_m[i]] = fsolve(lambda x: tmpfun(x[0], x[1], x[2]), [0.09, 0.25, 0.9])
In [13]: # incidence, screening and proportion of incident infections asymptomatic in women
        inc_f = np.zeros(n_sample)
        scr_f = np.zeros(n_sample)
        p_asymp_f = np.zeros(n_sample)
        for i in xrange(n_sample):
            def tmpfun(inc, scr, p_asymp):
                [tr, dr] = test_diag_fun(
                    array([
                        inc,
                        scr.
                        1-p_asymp, # proportion of incident infections which are symptomatic
                        sc_f[i], # rate of self-clear
                        att_symp[i],
                       p_true_pos_f[i],
                       p_false_pos_f[i]
                       ]))
                prev = dyn_fun(
                    inc*p_asymp,
                    sc_f[i] + scr*p_true_pos_f[i],
                    inc*(1-p_asymp),
                    scr*p_true_pos_f[i] + att_symp[i]*p_true_pos_f[i]
                return (tr - test_rate_f_15_24[i],
                        dr - diag_rate_f_15_24[i],
                       prev - prev_f[i])
            [inc_f[i], scr_f[i], p_asymp_f[i]] = fsolve(lambda x: tmpfun(x[0], x[1], x[2]), [0.09, 0.25, 0.9])
In [15]: h=plt.hist(p_asymp_f, bins=50, histtype='step', normed=True, color='r')
        h=plt.hist(p_asymp_m, bins=50, histtype='step', normed=True, color='b')
        plt.xlabel('Proportion of incident infections which are asymptomatic')
        print 'Median (central 95% credible interval) for proportion asymptomatic in men: \n \t', \
            percentile(p_asymp_m, 50), percentile(p_asymp_m, (2.5,97.5))
        print 'Median (central 95% credible interval) for proportion asymptomatic in women:\n \t', \
            percentile(p_asymp_f, 50), percentile(p_asymp_f, (2.5,97.5))
Median (central 95% credible interval) for proportion asymptomatic in men:
          0.509889142897 [0.26393408139570879, 0.75872661515902395]
Median (central 95% credible interval) for proportion asymptomatic in women:
          0.616465413009 [0.46763845173602908, 0.75205517865264992]
```

1.3 Estimating national prevalence

The sampled parameter values are now used to infer prevalence in men and women in different age groups.

```
In [15]: from scipy.optimize import fsolve
```

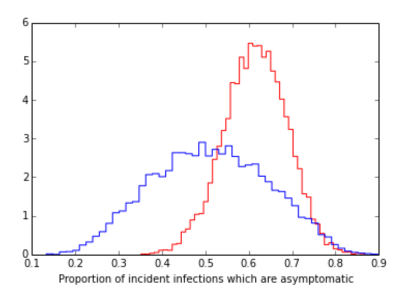


Figure 3: Samples for the proportion of incident infections which are asymptomatic in men (blue) and women (red), calibrated to Natsal-3 prevalence estimates in 16-24-year-olds.

```
In [16]: # men first...
         prev_m_15_19 = np.zeros(n_sample)
         inc_m_15_19 = np.zeros(n_sample)
scr_m_15_19 = np.zeros(n_sample)
         for i in xrange(n_sample):
              [inc_m_15_19[i], scr_m_15_19[i]] = fsolve(lambda x: test_diag_fun(concatenate([
                              x, array([
                                       1-p_asymp_m[i], # proportion of incident infections which are symptomatic
                                       sc_m[i], # rate of self-clear
                                       att_symp[i],
                                       p_true_pos_m[i],
                                       p_false_pos_m[i]
                                  ])])) - array([test_rate_m_15_19[i],diag_rate_m_15_19[i]]), [0.09, 0.25])
             prev_m_15_19[i] = dyn_fun(
                 inc_m_15_19[i]*p_asymp_m[i],
                 sc_m[i] + scr_m_15_19[i]*p_true_pos_m[i],
                 inc_m_15_19[i]*(1-p_asymp_m[i]),
                 scr_m_15_19[i]*p_true_pos_m[i] + att_symp[i]*p_true_pos_m[i]
             )
In [17]: prev_m_20_24 = np.zeros(n_sample)
         inc_m_20_24 = np.zeros(n_sample)
         scr_m_20_24 = np.zeros(n_sample)
         for i in xrange(n_sample):
             [inc_m_20_24[i], scr_m_20_24[i]] = fsolve(lambda x: test_diag_fun(concatenate([
                                       1-p\_asymp\_m[i] \text{, \# proportion of incident infections which are symptomatic} \\
                                       sc_m[i], # rate of self-clear
                                       att_symp[i],
                                       p_true_pos_m[i],
                                       p_false_pos_m[i]
                                  ])])) - array([test_rate_m_20_24[i],diag_rate_m_20_24[i]]), [0.09, 0.25])
             prev_m_20_24[i] = dyn_fun(
                 inc_m_20_24[i]*p_asymp_m[i],
                 sc_m[i] + scr_m_20_24[i]*p_true_pos_m[i],
                 inc_m_20_24[i]*(1-p_asymp_m[i]),
                 att_symp[i]*p_true_pos_m[i]
```

```
In [18]: # ... then women
        prev_f_15_19 = np.zeros(n_sample)
         inc_f_{15_19} = np.zeros(n_sample)
         scr_f_15_19 = np.zeros(n_sample)
        for i in xrange(n_sample):
             [inc_f_15_19[i], scr_f_15_19[i]] = fsolve(lambda x: test_diag_fun(concatenate([
                             x, array([
                                     1-p_asymp_f[i], # proportion of incident infections which are symptomatic
                                     sc_f[i], # rate of self-clear
                                     att_symp[i],
                                     p_true_pos_f[i],
                                     p_false_pos_f[i]
                                 ])])) - array([test_rate_f_15_19[i],diag_rate_f_15_19[i]]), [0.03, 0.44])
             prev_f_15_19[i] = dyn_fun(
                 inc_f_15_19[i]*p_asymp_f[i],
                 sc_f[i] + scr_f_15_19[i]*p_true_pos_f[i],
                 inc_f_15_19[i]*(1-p_asymp_f[i]),
                 scr_f_15_19[i]*p_true_pos_f[i] + att_symp[i]*p_true_pos_f[i]
In [19]: prev_f_20_24 = np.zeros(n_sample)
         inc_f_20_24 = np.zeros(n_sample)
         scr_f_20_24 = np.zeros(n_sample)
         for i in xrange(n_sample):
             [inc_f_20_24[i], scr_f_20_24[i]] = fsolve(lambda x: test_diag_fun(concatenate([
                             x, array([
                                     1-p_asymp_f[i], # proportion of incident infections which are symptomatic
                                     sc_f[i], # rate of self-clear
                                     att_symp[i],
                                     p_true_pos_f[i];
                                     p_false_pos_f[i]
                                 ])])) - array([test_rate_f_20_24[i],diag_rate_f_20_24[i]]), [0.03, 0.44])
             prev_f_20_24[i] = dyn_fun(
                 inc_f_20_24[i]*p_asymp_f[i],
                 sc_f[i] + scr_f_20_24[i]*p_true_pos_f[i],
                 inc_f_{20_24}[i]*(1-p_asymp_f[i]),
                 scr_f_20_24[i]*p_true_pos_f[i] + att_symp[i]*p_true_pos_f[i]
In [20]: # ...and now plot sampled prevalence by age group
         fig = plt.figure(figsize = (10,10))
         ax1 = fig.add_subplot(221)
        h_2012_m_15_19 = ax1.hist(
            prev_m_15_19, bins=20, normed=true, histtype='step', color='cyan', label='15-19 years')
        h_2012_m_20_24 = ax1.hist(
            prev_m_20_24, bins=20, normed=true, histtype='step', color='blue', label='20-24 years')
         ax1.errorbar(0.001, 25, xerr=[[0],[0.022-0.001]], ecolor='cyan', capsize=10)
         ax1.errorbar(0.022, 30, xerr=[[0],[0.052-0.022]], ecolor='blue', capsize=10)
         ax1.annotate('18-19 years', [0.001, 25], color='0.5')
        ax1.annotate('20-24 years', [0.022, 30], color='0.5')
        ax1.set_xlabel('Prevalence')
        ax1.set_xlim(0,0.1)
        ax1.set_ylim(0,115)
        ax1.set_title('Sexually active men')
        ax1.legend()
        ax2 = fig.add_subplot(222)
        h_2012_f_15_19 = ax2.hist(
             prev_f_15_19, bins=20, normed=true, histtype='step', color='fuchsia', label='15-19 years')
        h_2012_f_20_24 = ax2.hist(
             prev_f_20_24, bins=20, normed=true, histtype='step', color='r', label='20-24 years')
         ax2.errorbar(0.009, 20, xerr=[[0],[0.058-0.009]], ecolor='fuchsia', capsize=10)
         ax2.errorbar(0.025, 25, xerr=[[0],[0.086-0.025]], ecolor='fuchsia', capsize=10)
         ax2.errorbar(0.017, 30, xerr=[[0],[0.042-0.017]], ecolor='r', capsize=10)
```

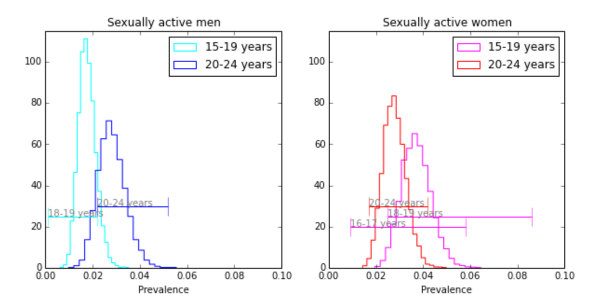


Figure 4: Sampled chlamydia prevalence in men (left) and women (right), by age group. Stepped histograms show samples. Horizontal bars give 95% confidence intervals for prevalence in comparable age groups, estimated from Natsal-3.

```
ax2.annotate('16-17 years', [0.009, 20], color='0.5')
        ax2.annotate('18-19 years', [0.025, 25], color='0.5')
        ax2.annotate('20-24 years', [0.017, 30], color='0.5')
        ax2.set_xlabel('Prevalence')
        ax2.set_xlim(0,0.1)
        ax2.set_vlim(0,115)
        ax2.set_title('Sexually active women')
        ax2.legend()
        print 'Central 95% credible interval for sexually active men, 15-19 years: \n \t', \
            percentile(prev_m_15_19, (2.5, 97.5))
        print 'Central 95% credible interval for sexually active men, 20-24 years: \n \t', \
            percentile(prev_m_20_24, (2.5, 97.5))
        print 'Central 95% credible interval for sexually active women, 15-19 years: \n \t', \
           percentile(prev_f_15_19, (2.5, 97.5))
        print 'Central 95% credible interval for sexually active women, 20-24 years: \n \t', \
            percentile(prev_f_20_24, (2.5, 97.5))
Central 95% credible interval for sexually active men, 15-19 years:
          [0.011220320772663614, 0.025540614007783301]
Central 95% credible interval for sexually active men, 20-24 years:
          [0.017855916166134259, 0.040426434826233246]
Central 95% credible interval for sexually active women, 15-19 years:
          [0.02590103035920896, 0.050182593529829664]
Central 95% credible interval for sexually active women, 20-24 years:
          [0.019236585373852842, 0.038203370665675189]
```

In these plots, step histograms show the sampled values for prevalence in men and women, by age group. The horizontal bars give 95% confidence intervals for prevalence in comparable age groups, estimated from Natsal-3. They show the agreement between our surveillance-based method and the population-based survey.

1.4 Symptomatic and asymptomatic diagnoses

Although the data does not report the number of diagnoses that were in symptomatic and asymptomatic cases, we can propose different possible numbers of symptomatic and asymptomatic diagnoses and examine the inferences which would have followed in each case.

```
In [21]: # men first...
         prev_m = np.zeros(n_sample)
         inc_m = np.zeros(n_sample)
         scr_m = np.zeros(n_sample)
         p_symp_m = np.zeros(n_sample)
         # there were 48387 diagnoses in men aged 15-24
         # don't allow all symptomatic or all asymptomatic - messes with gamma distributions
         sample_symp_m = ceil(48386*rs.uniform(size = n_sample))
         diag_rate_symp_m_15_24 = rs.gamma(sample_symp_m, 1, size=n_sample)/pop_active_m_15_24
         sample_asymp_m = 48387 - sample_symp_m
         diag_rate_asymp_m_15_24 = rs.gamma(sample_asymp_m, 1, size=n_sample)/pop_active_m_15_24
         for i in xrange(n_sample):
             [inc_m[i], scr_m[i], p_symp_m[i]] = fsolve(lambda x: test_diag_sym_asym_fun(concatenate([
                              x, array([
                                      sc_m[i], # rate of self-clear
                                      att_symp[i],
                                      p_true_pos_m[i]
                                      p_false_pos_m[i]
                                  ])])) - \
                                  arrav([
                                      test_rate_m_15_24[i],
                                      diag_rate_symp_m_15_24[i],
                                      diag_rate_asymp_m_15_24[i]
                                      ]),
                                  [0.01, 0.3, 0.21])
             prev_m[i] = dyn_fun(
                 inc_m[i]*(1-p_symp_m[i]),
                 sc_m[i] + scr_m[i]*p_true_pos_m[i],
                 inc_m[i]*p_symp_m[i],
                 sc_m[i] + scr_m[i]*p_true_pos_m[i] + att_symp[i]*p_true_pos_m[i])
In [22]: # ...then women
         prev_f = np.zeros(n_sample)
         inc_f = np.zeros(n_sample)
         scr_f = np.zeros(n_sample)
         p_symp_f = np.zeros(n_sample)
         # there were 88101 diagnoses in women aged 15-24
         # don't allow all symptomatic or all asymptomatic - messes with gamma distributions
         sample_symp_f = ceil(88100*rs.uniform(size = n_sample))
         diag_rate_symp_f_15_24 = rs.gamma(sample_symp_f, 1, size=n_sample)/pop_active_f_15_24
         sample_asymp_f = 88101 - sample_symp_f
         \label{limits}  \mbox{diag\_rate\_asymp\_f\_15\_24 = rs.gamma(sample\_asymp\_f, 1, size=n\_sample)/pop\_active\_f\_15\_24} \\
         for i in xrange(n_sample):
             [inc_f[i], scr_f[i], p_symp_f[i]] = fsolve(lambda x: test_diag_sym_asym_fun(concatenate([
                              x, array([
                                      sc_f[i], # rate of self-clear
                                      att_symp[i],
                                      p_true_pos_f[i]
                                      p_false_pos_f[i]
                                  ])])) - \
                                  array([
                                      test_rate_f_15_24[i],
                                      diag_rate_symp_f_15_24[i],
                                      diag_rate_asymp_f_15_24[i]
                                      ]),
                                  [0.01, 0.3, 0.21])
             prev_f[i] = dyn_fun(
```

```
inc_f[i] * (1-p_symp_f[i]),
                 sc_f[i] + scr_f[i]*p_true_pos_f[i],
                 inc_f[i]*p_symp_f[i],
                 sc_f[i] + scr_f[i]*p_true_pos_f[i] + att_symp[i]*p_true_pos_f[i])
In [23]: fig = plt.figure(figsize = (10,12))
         xtk_m = [0, 10000, 20000, 30000, 40000] # x-axis ticks for men
         xtk_f = [0, 20000, 40000, 60000, 80000] # x-axis ticks for women
         ax1 = fig.add_subplot(421)
         ax1.plot(100*sample_symp_m/48387, prev_m, ".", alpha = 0.1)
         ax1.fill_between([0,50000], 0.015, 0.034, facecolor="b", alpha=0.3)
         ax1.plot([60,60],[0,1],"--b")
         ax1.plot([80,80],[0,1],"--b")
         ax1.set_xlim([0,100])
         ax1.set_ylim([0,0.1])
         ax1.set_ylabel("Prevalence")
         ax1.set_title("Sexually active men, 15-24 years")
         ax2 = fig.add_subplot(422)
         ax2.plot(100*sample_symp_f/88101, prev_f, ".r", alpha = 0.1)
         ax2.fill_between([0,100000], 0.022, 0.043, facecolor="r", alpha=0.3)
         ax2.plot([45,45],[0,1],"--r")
         ax2.plot([70,70],[0,1],"--r")
         ax2.set_xlim([0,100])
         ax2.set_ylim([0,0.1])
         ax2.set_title("Sexually active women, 15-24 years")
         ax3 = fig.add_subplot(423)
         ax3.plot(100*sample_symp_m/48387, inc_m, ".", alpha = 0.1)
         ax3.plot([60,60],[0,1.2],"--b")
         ax3.plot([80,80],[0,1.2],"--b")
         ax3.set_xlim([0,100])
         ax3.set_ylim([0,0.2])
         ax3.set_ylabel("Incidence")
         ax4 = fig.add_subplot(424)
         ax4.plot(100*sample_symp_f/88101, inc_f, ".r", alpha = 0.1)
         ax4.plot([45,45],[0,1.2],"--r")
         ax4.plot([70,70],[0,1.2],"--r")
         ax4.set_xlim([0,100])
         ax4.set_ylim([0,0.2])
         ax5 = fig.add_subplot(425)
         ax5.plot(100*sample_symp_m/48387, scr_m, ".", alpha = 0.1)
         ax5.plot([60,60],[0,1],"--b")
         ax5.plot([80,80],[0,1],"--b")
         ax5.set_xlim([0,100])
         ax5.set_ylim([0,0.5])
         ax5.set_ylabel("Screening")
         ax6 = fig.add_subplot(426)
         ax6.plot(100*sample_symp_f/88101, scr_f, ".r", alpha = 0.1)
         ax6.plot([45,45],[0,1],"--r")
         ax6.plot([70,70],[0,1],"--r")
         ax6.set_xlim([0,100])
         ax6.set_ylim([0,0.5])
         ax7 = fig.add_subplot(427)
         ax7.plot(100*sample_symp_m/48387, p_symp_m, ".", alpha = 0.1)
         ax7.plot([60,60],[0,1],"--b")
         ax7.plot([80,80],[0,1],"--b")
         ax7.plot([0,100],[0.24,0.24],"--b")
         ax7.plot([0,100],[0.74,0.74],"--b")
         ax7.set_xlim([0,100])
         ax7.set_ylim([0,1])
         ax7.set_xlabel("Proportion of diagnoses symptomatic (%)")
         ax7.set_ylabel("Proportion of incident infections symptomatic")
```

```
ax8 = fig.add_subplot(428)

ax8.plot(100*sample_symp_f/88101, p_symp_f, ".r", alpha = 0.1)

ax8.plot([45,45],[0,1],"--r")

ax8.plot([70,70],[0,1],"--r")

ax8.plot([0,100],[0.25,0.25],"--r")

ax8.plot([0,100],[0.53,0.53],"--r")

ax8.set_xlim([0,100])

ax8.set_ylim([0,1])

Out[23]: (0, 1)
```

The dashed lines are intended as a guide to the eye, to indicate scenarios roughly compatible with the Natsal-3 prevalence estimates. The observed chlamydia prevalence in Natsal-3 would be consistent with around 60-80% of diagnoses in men and 45-70% in women being symptomatic.

```
In [24]: # plot top pair only, for figure in paper
        fig = plt.figure(figsize = (10,3))
        xtk_m = [0, 10000, 20000, 30000, 40000] # x-axis ticks for men
        xtk_f = [0, 20000, 40000, 60000, 80000] # x-axis ticks for women
        ax1 = fig.add_subplot(121)
        ax1.plot(100*sample_symp_m/48387, prev_m, '.', alpha = 0.1)
        ax1.fill_between([0,50000], 0.015, 0.034, facecolor='b', alpha=0.3)
        ax1.plot([60,60],[0,1],'--b')
        ax1.plot([80,80],[0,1],'--b')
        ax1.set_xlim([0,100])
        ax1.set_ylim([0,0.1])
        ax1.set_xlabel('Proportion of diagnoses symptomatic (%)')
        ax1.set_ylabel('Prevalence')
        ax1.set_title('Sexually active men, 15-24 years')
        ax2 = fig.add_subplot(122)
        ax2.plot(100*sample_symp_f/88101, prev_f, '.r', alpha = 0.1)
        ax2.fill_between([0,100000], 0.022, 0.043, facecolor='r', alpha=0.3)
        ax2.plot([45,45],[0,1],'--r')
        ax2.plot([70,70],[0,1],'--r')
        ax2.set_xlim([0,100])
        ax2.set_ylim([0,0.1])
        ax2.set_xlabel('Proportion of diagnoses symptomatic (%)')
         ax2.set_title('Sexually active women, 15-24 years')
Out[24]: <matplotlib.text.Text at 0x111d06050>
```

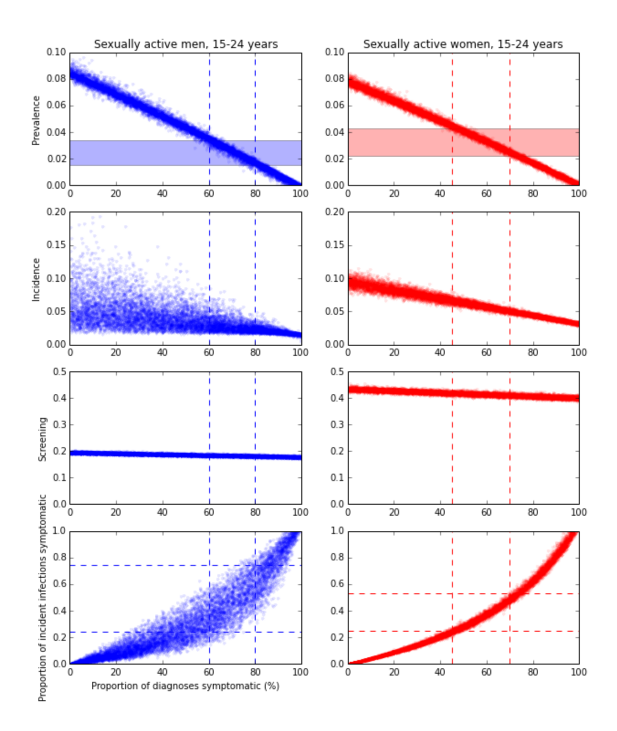


Figure 5: Samples for prevalence, incidence, screening rate and proportion of infections which are symptomatic, assuming different proportions of diagnoses made as a result of symptomms. The dashed lines are intended as a guide to the eye, to indicate scenarios roughly compatible with the Natsal-3 prevalence estimates.

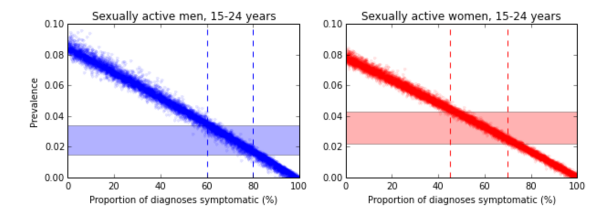


Figure 6: The upper two panels from the previous figure.