Chlamydia 2000

Estimating Reinfection Intervals for

*Chlamydia trachomatis*

based on Routine Data Collection

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Declaration

I, Florian Burckhardt, declare the following dissertation to be my own work and entirely composed by myself.
Acknowledgements

I would like to thank the following people for their cooperation and help in writing this dissertation:

My tutor Pamela Warner, for her guidance, advice and time spent on discussions. Her patience would do a Zen-Master proud.

Sheena Sutherland for allowing me to use her data and for her helpful suggestions.

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John Young and Gordon Scott for helpful information on study related issues.

Moral support: Coco for sending Comics&Chocolate, Markus in General, Kaffe Politik for their carrot cake, Sid Meier for his Game Alpha Centauri & Friends

I dedicate this dissertation to my granddad Rudolf Külbel.
A lord of ancient China once asked his physician, a member of a family of healers, which of them was the most skilled in the art. The physician, whose reputation was such that his name became synonymous with medical science in China, replied:

"My eldest brother sees the spirit of sickness and removes it before it takes shape, so his name never gets out of the house.

"My elder brother cures sickness when it is still extremely minute, so his name does not get out of the neighbourhood.

"As for me, I puncture veins, prescribe potions, and massage skin, so from time to time my name gets out and is heard among the lords.

Chinese Tale
Abstract

*Chlamydia trachomatis* is the most common bacterial sexually transmitted disease in Scotland and the rest of the UK. Its sequelae include pelvic inflammatory disease, ectopic pregnancy, infertility and arthritis and these are more likely if reinfection occurs.

The costs to the healthcare system are estimated at £50 million a year and increased resources have been directed towards piloting a national screening program in the UK. Due to the nature of the disease, reinfection is common and knowledge of the time between subsequent infections is important for retest intervals in screening programs. Despite numerous studies on reinfection with *Chlamydia*, the actual reinfection interval for a British GUM clinic population is not known.

This study analyses routine data on *Chlamydia* tests collected retrospectively from January 1992 until May 2000 by the Medical Microbiology Laboratory of the Royal Infirmary Edinburgh GUM clinic. A total of 47,587 tests made on 34,754 patients were analysed with survival methods to estimate risk-group specific reinfection intervals and to identify the importance of factors available for analysis that may be determinants of reinfection. Variables were examined to ensure the assumptions underlying the analyses were met.

The process of data cleaning, analysis and the rationale behind it are described in detail because of their importance in studies using routinely collected data and to enable similar studies on routine data of other GUM clinics. Results are discussed and areas for future research identified.
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I. Introduction

Epidemiology can be seen as the study of the pattern of disease through time, place and population. It seeks to uncover the hidden links and causations of ill-health on a population rather than a physiological, individual level. Epidemiological studies look at the associations between risk factors (exposures) and disease outcomes. They can try to infer causations from data in order to create hypotheses of why people get a disease. Alternatively, if the aetiology of a disease is already known in detail, knowledge of who is more likely to get the disease is essential for cost efficient medical and educational support. This is even more important for risk factors that lie beyond an individual's control such as age or ethnicity. It is an ethical obligation to be as efficient in delivering health care as possible, since resources wasted unnecessarily are not available to others.

Sexually transmitted diseases (STDs) are a major burden of disease worldwide and bring great suffering. They can lead to severe medical consequences such as infertility for both, men and women, adverse pregnancy outcomes or even death and cause a high social and economic burden. STDs share overlapping epidemiologies with similar modes of transmissions and symptoms. Any insight into the underlying disease patterns of one STD could possibly be transferrable to others. STDs are more controlled by behaviour than by physiological constitution. Precise knowledge of risk groups allows targeted prevention strategies such as specialised health education or better access to screening programs.

*Chlamydia trachomatis* is causing the majority of sexually transmitted bacterial infections throughout the world. With efficient diagnostic tests and treatment for the disease at disposal, *Chlamydia* infections challenge the non-medical aspects of Public Health, such as identifying and targeting risk groups or providing education and access to healthcare.

Despite lots of dedicated research, one of the important open questions regarding *Chlamydia* is that of the reinfection interval. Ideally, one would like to be able to define an interval based on selected personal characteristics of an individual. Multivariate statistical methods can help to find the "right" set of characteristics in a study population. However, it has to be checked carefully whether findings can be generalised to other settings. For an adequate Public Health response to *Chlamydia*, one would have to consider not only reinfection intervals, but also qualitative information of group specific seriousness of sequelae and access to healthcare infrastructure. Health economic and resource management implications would have to be considered, too.
With the widespread use of modern data processing in a lot of health care settings, routinely collected data can be accessed quickly without time consuming compilation of written records. Multicentre databases are making an increasing contribution to medical understanding as they allow one to tap into a rich seam of epidemiological data for retrospective studies.

This study analyses routine data on Chlamydia tests collected retrospectively from January 1992 until May 2000 by the Medical Microbiology Laboratory of the Royal Infirmary Edinburgh GUM clinic. A total of 47,587 tests made on 34,754 patients are analysed with survival methods to estimate risk-group specific reinfection intervals and to identify determinants of reinfection. The large time interval makes the study sample one of the largest ever on Chlamydia in the UK and one of the largest worldwide involving both, men and women.

The process of data cleaning, analysis and the rationale behind it are described in detail because of their importance in studies using routinely collected data and to enable similar studies on routine data of other GUM clinics.

Outline

Chapter II will cover epidemiological issues by reviewing the current literature on Chlamydia. It will also give a brief microbiological background.

Chapter III will describe the study design and give details on the routine data collection process.

Chapter IV will introduce the statistical methods used and describe the analyses made.

Chapter V will provide more detailed information on data storage and retrieval, which would otherwise have obstructed the reading flow.

Chapter VI will report the results in tables and figures.

Chapter VII will discuss the results of this study, reflect on its implications and make recommendations.

The appendix contains a list of abbreviations used in this dissertation.
II. Review of the Literature

Overview

*Chlamydia trachomatis* is the most common bacterial sexually transmitted disease (STD) in Scotland (ISD, 2000) and the rest of the UK (Stephenson, 1998). The infection is asymptomatic in 50% of men and 70% of women (CMO, 1997) and can thus be passed on quite readily before any preventative or curative measures are taken.

*Chlamydia* infections have major medical, social and economic consequences. Pelvic inflammatory disease (PID), ectopic pregnancy, tubal factor infertility and epididymitis, proctitis and arthritis (Paavonen *et al*, 1996) are all extremely costly sequelae to the healthcare system with conservative calculations being estimated at £50 million per year (Stephenson, 1998). Women are particularly affected with further adverse outcomes including chronic pelvic pain, premature rupture of membranes during pregnancy, low birth weight of infants, still birth and early pregnancy loss. In neonates of infected mothers, *Chlamydia* conjunctivitis, trachoma (hence the name) and pneumonitis may develop (Genc *et al*, 1996). It is also estimated that 6 million people lost their eyesight because of *Chlamydia* infections (Kayser *et al*, 1992). In the tropics, C. trachomatis is responsible for lymphogranuloma venerum.

*Chlamydia* infections are also linked to an increased susceptibility to HIV, probably due to the inflammatory response that leads to a higher concentration of HIV-host cells (Royce *et al*, 1997). In what follows, only sexually transmitted *Chlamydia* infections are considered.

In addition to any economic cost, the psychological burden for an individual suffering from infertility, chronic PID or having survived an ectopic pregnancy will be severe. As a result of C. trachomatis infection, in the UK alone each year about 74,000 women will suffer from PID, 30,000 couples will seek fertility treatment and 3,000 ectopic pregnancies will occur, 120 of which will lead to death of the mother (Taylor-Robinson, 1994). There is clearly a strong public health interest in reducing infection and reinfection with *Chlamydia*, which has led to the launching of a national screening study pilot in the UK (Bower, 1998, Department of Health, 2000)

One of the key issues for future research pointed out by the Chief Medical Officers’ (CMO) expert advisory group on *Chlamydia* concerns optimum screening intervals (CMO, 1997). Even the most recent report on a national screening pilot study (Tobin *et al*, 2000) identifies this as a crucial question, that remains to be answered. Screening intervals will depend on reinfection probabilities and intervals, access
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to risk groups, severity of sequelae, existing health infrastructure and resources available. Methods for estimation of reinfection intervals and reporting them is the focus for this dissertation.

Microbiological Background

*Chlamydia trachomatis* belongs to the *Chlamydiaceae*, a group of obligate intracellular bacterial parasites (Kayser *et al*, 1992). For the remainder of the text, “*Chlamydia*” (genus) will refer to *Chlamydia trachomatis* (species) unless stated otherwise. *Chlamydiaceae* differ from other bacteria by going through a special reproductive cycle with two distinct morphological stages, the infectious elementary bodies (EB) and the reproductive reticulate bodies (RB).

An elementary body is about 300 nm wide, dense, spherical and with a rigid cell wall especially adapted to survive outside a host cell. It also contains the necessary receptors to dock onto the outside of mucosal host cells and to trigger its own phagocytosis, thus conveying infectivity. Once inside the cellular compartments of a mucosal cell, EBs change to become the larger (1000 nm), less dense and non-infectious RBs that grow through cellular division inside their host cell and drain its resources.

Subsequently, some RBs change back to become EBs. Upon lysis of the host cell, both RBs and EBs get released and the EBs continue the infectious cycle. One cycle from docking on the host to lysis of the host takes about 48h (Kayser *et al*, 1992; appendix, fig. 1).

*C. trachomatis*, like all *Chlamydiaceae*, exists in a wide range of different serotypes, which are responsible for different sequelae. Host acquired immunity against one serotype is partial since it does not protect against a different serotype, thus making subsequent infections possible.

A host's immune system has, simply put, two main strategies, humoral (non-cellular) and cellular defence. Humoral defence consists mainly of different types of antibodies dissolved in blood plasma, ready to attack and immobilise any pathogen they encounter and able to call in “help” from lymphocytes. Cellular defence consists of specialized lymphocytes such as natural killer cells that can recognise and kill “invaded” body cells. Being an intracellular parasite, *Chlamydia* basically evades humoral immune defence and cellular defence only can be effective. There is growing evidence now for reinfections being associated with chronic inflammation and increasing the risk for ectopic pregnancy through an excessive inflammatory response with a subsequent scarring of tissue, which causes tubal blockage (Hillis *et al*, 1997, Rasmussen *et al*, 1997, Patton *et al*, 1989).

Further issues surrounding reinfection will be discussed in greater detail later.
Diagnosis of *Chlamydia*

A lot of veneral infections share overlapping symptoms and can also be present simultaneously (Fortenberry *et al.*, 1999), so diagnosis of infection is a key step. For *Chlamydia*, proof of live culture used to be the method of choice because of its high specificity. With the discovery of monoclonal antibodies, immunofluorescent methods (direct fluorescent antibodies, enzyme immunoassays) were also used in detecting *Chlamydia* (CEG, 1999). The antibodies targeted an outer membrane protein of EBs that is shared by all serotypes. However, this still required invasive sampling and RBs could escape detection. The advent of DNA-amplification made it possible to amplify specific *Chlamydia*-only sequences, even with very diluted specimen such as a patient's urine. Studies have shown that the new tests have a higher sensitivity and specificity than previous tests (Quinn *et al.*, 1996, Young *et al.*, 1998).

It is obvious that a test should have a high sensitivity to pick up positives, but it also should be specific, otherwise false positive results would cause unnecessary worries for the individuals concerned (CMO, 1997). This is more likely to happen where the prevalence of the condition in a population is low. Sensitivity is about 75% – 100%, specificity >99% when used on non-invasive samples like first void urine (FVU) (CEG, 1999). The test can also detect *C. trachomatis* infection when organisms are in very low numbers, which is important for early diagnosis. Testing is also less dependent on sampling and transportation techniques (Stary, 1997), so even home sampling of FVU might be an option.

Men especially benefit from the new non-invasive sampling, as the previous method involved rather painful urethral swabs. Indeed, since introduction of the new testing method the total number of men and also the relative proportion of men testing positive has increased, because more partners of positive women agreed to get tested (Dr. Sheena Sutherland, affil.).

A particular diagnostic problem is inherently connected with the high sensitivity of amplification assays. Based on DNA amplification and able to detect minute amounts of it, undegraded DNA from dead or non viable bacteria could give a false positive result if tested within 3 weeks of initial treatment. Therefore, a test of cure (TOC) has to be made 3 weeks after treatment (CEG, 1999).

**Treatment**

*As a bacterium, Chlamydia is vulnerable to antibiotics. Antibiotics of choice are tetracyclines and macrolides. The infection is easily treated with either Doxycycline (100mg) or Erythromycin (500mg) for 7 days or Azithromycin (1000mg) given in a single dose (Martin *et al.*, 1992, CEG 1999). Azithromycin guarantees compliance, as doctors can observe patients taking the treatment, but it is almost 4 times more effective*
expensive (Stephenson, 1998). Unlike a lot of other antibiotics, Acithromycin is still under patent (Pfizer Pharmaceuticals), so its holder can dictate the price. This raises issues of patient compliance with treatment vs. cost of treatment, which have to be balanced carefully for different public health settings. CEG (1999) recommends Acithromycin for patients with erratic healthcare seeking behavior. There is evidence for Acithromycin having overall cost advantages, however, mainly because of its 100% compliance rate (Black et al., 2000).

*Chlamydia* is an intracellular parasite with an unusual life cycle, therefore genetic exchange of resistance plasmids with other bacteria will be extremely limited and no antibiotic resistance is known (Young et al., 1998). This is important for practical management, since concerns regarding non-compliance are limited to issues of cure of patient and reduction of infection pool. There is no danger of antibiotic resistance developing due to non-compliance.

Prevalence and Risk Factors

The majority of studies on *Chlamydia* were conducted on women only (table 1.1-2). The main reasons might be availability of routine data, accessibility of study population and severity of sequelae. Men are less likely than women to attend healthcare settings where screening would be feasible and their sequelae are less severe (Tobin et al., 2000). There is also a far more extensive reproductive health infrastructure available for women than for men, e.g. routine cervical cancer screening. It is now recommended practice to include testing for *Chlamydia* in all these health settings (SIGN, 2000).

In addition, the moral pressure on women regarding STDs is certainly higher than that on men. However, one must not ignore the contribution of men in spreading STDs and more research in that area would help to get a better overall picture on the pattern of disease (Pierpoint et al., 2000).

The exact prevalence of *C. trachomatis* is not known but numbers for women range from 3% to 11% (James, 1999, Oakeshott et al., 1995, Paavonen, 1997, SCIEH, 1999, Santer et al., 2000). However, it is very clear from routine data on sexually transmitted diseases (ISD, 2000, Simms et al., 1997) and from other studies (Stokes, 1997, Grun et al., 1997) that risk of infection is highly age dependent, with highest prevalence in teenage women and peak levels for men aged 25-34.

Within Scotland, Lothian accounts for almost a quarter of all cases (ISD, 2000). The number of positives in Scotland has increased by 20% annually since 1996 (ISD, 2000). Part of the increase can be attributed to the higher sensitivity of the new LCx test, since an increasing number of laboratories were shifting to the new amplification assays during the last years (SCIEH, 1999). In case of the Royal Infirmary data,
test methods changed from culture and immunofluorescence to LCx in summer 1998, leading to an immediate 1.5 increase in positive diagnoses.

Effects at population level are determined by the behaviours of individuals. In theory, sexual transmission can be prevented almost completely by using condoms. In real life on the other hand, behavioural risk factors and socioeconomic proxy measures are used to explain the observed differences in infection rates within a population: young age, ethnic group, low school leaving age, single status, not using barrier contraceptives, multiple sexual partners or a new partner in recent months are considered to be risk factors (table 1.1-2). However, some studies contradict the findings of others. In a study by Burstein et al (1998), common predictors such as prior STD-history, multiple or new partners and inconsistent condom use were, however, not able to identify a high-risk subset among adolescent females. Regarding hormonal contraception, one study reported a protective effect (Richey et al, 1999) and another found the opposite (CMO, 1997). This might be explained by different kinds of sexual relationships of women taking hormonal contraception: active family planning in a secure relationship combined with a low risk attitude vs. casual relationships with convenient pregnancy prevention.

Cultural differences between study settings will lead to different conclusions and recommendations. For example, ethnicity is used as a covariate in most of the studies (tab. 1.1-2). In most US studies, however, the ethnicity variable only accounts for “white”, “black” and “other” (Blythe et al, 1992, Hillis et al, 1994, Fortenberry et al, 1999, Richey et al, 1999), in UK studies it additionally differentiates “black Caribbean”, “black African”, “Asian” (Hughes et al, 2000, Shahmanesh et al, 2000). It is unclear to what extent findings for a racial subgroup as risk factor can be generalised to other settings.

Summing up, for women, young age seems to be the most robust predictor for increased risk of Chlamydial infection. With regard to men, there has been too little data to establish robust predictors of increased risk.
Estimating Reinfetion Intervals for *Chlamydia trachomatis*

Table 1.1: Studies on risk factors for *Chlamydia* infection.

<table>
<thead>
<tr>
<th>Author</th>
<th>study type</th>
<th>Sex</th>
<th>Age</th>
<th>Country</th>
<th>L(P)CR</th>
<th>Risk Factors, other findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMO Expert Advisory Group, 1997</td>
<td>summary</td>
<td>both</td>
<td>all</td>
<td>UK</td>
<td>various</td>
<td>young age, ethnic group, single status, oral contraceptives, new sexual partners within last 3 months, no previous births, low school leaving age</td>
</tr>
<tr>
<td>Hughes, 2000</td>
<td>cross sectional</td>
<td>both</td>
<td>all</td>
<td>UK</td>
<td>various</td>
<td>black ethnic minority, teenagers, multiple partners</td>
</tr>
<tr>
<td>Mosure, 1996</td>
<td>retrosp.</td>
<td>women</td>
<td>15-19</td>
<td>US</td>
<td>no</td>
<td>cervicitis, friable cervix, multiple/ new/ symptomatic sex partners; study population: more than one visit to family planning clinic</td>
</tr>
<tr>
<td>Pierpoint, 2000</td>
<td>cross sectional</td>
<td>men</td>
<td>only</td>
<td>UK</td>
<td>yes</td>
<td>low response rate (51%), prevalence 1.9%, highest in men &gt;30, screening women and contact tracing male partners may be efficient for <em>Chlamydia</em> control</td>
</tr>
<tr>
<td>Shahmanesh, 2000</td>
<td>cross sectional</td>
<td>both</td>
<td>all</td>
<td>UK</td>
<td>(no) *</td>
<td>within large urban centres, <em>Chlamydia</em> infections occur in core areas</td>
</tr>
<tr>
<td>Simms, 1997</td>
<td>retrosp.</td>
<td>both</td>
<td>all</td>
<td>UK</td>
<td>no</td>
<td>16-19 year old, particularly women; high levels of asymptomatics</td>
</tr>
<tr>
<td>Winter, 2000</td>
<td>retrosp.</td>
<td>both</td>
<td>15-64</td>
<td>UK</td>
<td>no</td>
<td>men: ethnic group, women: young age, interactions between ethnic group and age for both sexes and ethnic group and level of deprivation for men; ecological study</td>
</tr>
</tbody>
</table>

Studies on risk factors for reinfection are inconclusive (table 1.2). Young age, multiple/ new partners, presence of other STDs and ethnic group increase the risk of reinfection in studies with women only (Fortenberry *et al*, 1999, Hillis *et al*, 1998, Hillis *et al*, 1994) but not in others which also include men (Miller *et al*, 1998, Richey *et al*, 1999). Reinfection rate ranged between 17% and 54% and reinfection intervals, where given, between 6 months and 1 year (Kjaer *et al*, 2000, Blythe *et al*, 1992, Fortenberry *et al*, 1999).

Although *Chlamydia* is the most widespread STD in the western world, one still needs either a high-risk group or a very large sample to detect reinfection events. Most studies therefore take either large datasets from GUM clinics, family planning clinics or other health care setting (Hillis *et al*, 1994, Miller *et al*, 1998, Richey *et al*, 1999) or enroll adolescent women, a high risk group, for a prospective cohort study (Blythe *et al*, 1992, Fortenberry *et al*, 1999). Pimenta *et al* (2000) have analysed reinfection rates in England and found far lower rates (3.6%-9.4%) than those reported from the US studies. Treatment success of initial infections is high (95%) and within rates of pharmacological treatment failure (Hillis *et al*, 1998). Therefore, a reinfection event will most likely come from a new or an untreated partner (Blythe *et al*, 1992). This points out the importance of consequent contact tracing and partner treatment, which will be discussed below.
Table 1.2: Studies on risk factors for reinfection with *Chlamydia*.

<table>
<thead>
<tr>
<th>Author</th>
<th>study type</th>
<th>Sex</th>
<th>Age</th>
<th>Country</th>
<th>L(P)CR</th>
<th>Risk Factors, other findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Burstein, 1998</td>
<td>prospective</td>
<td>women</td>
<td>12-19</td>
<td>US</td>
<td>yes</td>
<td>included risk factors (prior disease, multiple/new partners, inconsistent condom use) failed to identify a high risk subset, reinfection interval: 6.3 months</td>
</tr>
<tr>
<td>Blythe, 1992</td>
<td>prospective</td>
<td>women adolescent</td>
<td></td>
<td>US</td>
<td>no</td>
<td>38.4% reinfection, majority within 9 months, reinfections with same serovar frequent, suggesting relapse or reinfection from untreated partner</td>
</tr>
<tr>
<td>Fortenberry, 1999</td>
<td>prospective</td>
<td>women</td>
<td>15-19</td>
<td>US</td>
<td>no</td>
<td>ethnic group, gonorrhea as initial infection, multiple sex partners in previous 3 months, inconsistent condom use, 40% recurrence with at least one STD within one year</td>
</tr>
<tr>
<td>Hillis, 1998</td>
<td>prospective</td>
<td>women</td>
<td>all</td>
<td>US</td>
<td>yes</td>
<td>2-3 fold increased risk for: &lt;24 and white, multiple/new partners, untreated partner</td>
</tr>
<tr>
<td>Hillis, 1997</td>
<td>retrosp.</td>
<td>women</td>
<td>all</td>
<td>US</td>
<td>no</td>
<td>&lt;25 years, black, place of residence</td>
</tr>
<tr>
<td>Hillis, 1994</td>
<td>retrosp.</td>
<td>women</td>
<td>&lt;15 - 44</td>
<td>US</td>
<td>no</td>
<td>young age, ethnic group, area of residence, coinfection with gonorrhea, STD history; receiving care in a family-planning clinic protective; 54% (&lt;15) and 30% (15-19) reinfection within 5 years</td>
</tr>
<tr>
<td>Kissinger, 1998</td>
<td>prospective</td>
<td>women</td>
<td>14-39</td>
<td>US</td>
<td>no</td>
<td>annual recurrence rate lower for patient delivered partner medication (11.5%) compared to partner referral group (25.5%)</td>
</tr>
<tr>
<td>Kjær, 2000</td>
<td>prospective</td>
<td>both</td>
<td>&gt;18</td>
<td>DK</td>
<td>yes</td>
<td>presence of other STDs associated with higher risk of reinfection, cumulated incidence of recurrence within 24 weeks: 29%; home sampling promising method for retesting</td>
</tr>
<tr>
<td>Miller, 1998</td>
<td>retrosp.</td>
<td>women</td>
<td>all</td>
<td>US</td>
<td>no</td>
<td>young age, pregnancy, infection with other STDs not predictive for reinfection with <em>Chlamydia</em>, 17% reinfection</td>
</tr>
<tr>
<td>Pimenta, 2000</td>
<td>retrosp.</td>
<td>women</td>
<td>?</td>
<td>UK</td>
<td>?</td>
<td>3.6% overall reinfection rate per year, black Carribians, multiple partners, previous STD</td>
</tr>
<tr>
<td>Richey, 1999</td>
<td>retrosp.</td>
<td>both (few males)</td>
<td></td>
<td>US</td>
<td>no</td>
<td>reinfection risk independent of age, multiple/new partners or other STDs; reduced risk of reinfection associated with tubal litigation, hormonal/barrier contraception; number of visits to clinic protective</td>
</tr>
</tbody>
</table>

* inconsistently reported
Control Strategies

Screening and contact tracing are the key strategies discussed for STD control (Tobin et al, 2000). An infective agent with a large pool of asymptomatic carriers unaware of their condition can spread extensively into the population before preventative measures are taken. One strategy of detecting asymptomatics is opportunistic screening during routine health visits such as cervical smear tests or during special treatments like termination of pregnancy (SIGN, 2000, Santer et al, 2000). A pilot study for nationwide Chlamydia screening has been set up in Portsmouth and the Wirral and offers opportunistic screening for women aged 16-25 who attend GPs, family planning, termination of pregnancy, genitourinary medicine (GUM), colposcopy, gynaecology, or antenatal clinics (Department of Health, 2000).

On the other hand, infections can only occur in sex partners of an infected index case. Therefore, another strategy for detecting infection in asymptomatics is by following up sex partners of a positive index case. This is termed contact tracing, a vital part in STD management because, as the name implies, these infections are transmitted by having sex. The proportion of asymptomatic infections in sexual partners was about 60% in a Danish study (Kjaer et al, 2000). The best test and treatment efforts are foiled if the partner of an index case is not tested and treated as well, because a ping-pong-like effect would then lead to reciprocal infections between partners (Blythe et al, 1992). With high rates of contact tracing, however, it possible to lower prevalence close to eradication. Kretzschmar et al (1996) have modelled different strategies for STD management: mass screening, focal screening and contact tracing. In their simulations, they found that Chlamydia needed much higher rates of contact tracing than other STDs in order to achieve eradication.

Sweden has very high contact tracing rates and even has legislation in place that allows for police enforced testing of named partners (Tyden et al, 2000). In the Royal Infirmary health care setting, less than 30% of partners of women tested positive came forward for testing (Dr. Sheena Sutherland, affil.). The previously mentioned Danish study that detected a high rate of asymptomatics, offered home sampling of first void urine. This approach takes into account the social reality of STDs in that it provides anonymity and thus avoids stigmatisation. In addition, home sampling is very convenient. Sacrificing a small amount of sensitivity for a two to threefold increase in partner participation rate warrants careful consideration as a future option for contact tracing. The RIEGUM clinic has already secured funding for home sampling and plans to offer this option in the second half of 2001 (Dr. Gordon Scott, personal communication).
Modelling

Risk factor studies, both retrospective and prospective, are empirical and inductive. They try to infer from data the traits and characteristics that make an individual member of the study population more likely to get the disease in question. Provided the study population is representative, the findings can then be extrapolated to a larger population and help in making the appropriate healthcare management decisions to lower morbidity.

A different epistemological approach to gain insight into disease patterns is mathematical model building. During this rather deductive process, a model for the spread of disease within a population through time is formulated after conceptual reflection and theoretical inquiry. One advantage is that any assumption is made explicit and can be scrutinised carefully. The model is usually translated into a computer simulation program and fed with starting values from empirical observations.

Mathematical modelling of STDs is fairly new (Anderson et al, 1991, Diekmann et al, 2000) and draws on a variety of different disciplines, including the social sciences (Wasserman et al, 1994). A model can be deterministic (Renshaw, 1991) or stochastic (Kretzschmar et al, 1996) and have virtually any degree of complexity. The difficulty lies in reducing the complexity as much as possible while keeping it as close to reality as possible. A model must not require estimation of more parameters than can sensibly be derived from data (Garnett et al, 1996).

“Classic” deterministic models are often an extension of the old Lotka-Volterra predator-prey differential equations (Lotka, 1925, Volterra, 1926) to accommodate host-parasites relationships (Renshaw, 1991). Basically, infected and non-infected are seen as different compartments which are connected with each other and have different influx and efflux rates.

Stochastic models also have different compartments for infected and non-infected, but a transition matrix of probabilities replaces fixed (“deterministic”) exchange rates (Kretzschmar et al, 1996).

Models can be expanded to account for the complexity of social networks within a population by splitting the population into a high prevalence group, e.g. young, and a low prevalence group, e.g. old people (Kretzschmar et al, 1996, Aral et al, 1999). Including social networks would account for the fact that population dynamics does not merely consist of the sum of its individuals but includes the interactions between them as well. As Koopman et al (1999) argues, this would take into account the “network” plane of epidemiological data, i.e. the arrangement of and exchange between individuals, which is lost by merely looking at the “individual” plane of “classic” epidemiological studies with exposure and outcome variables per individual only.
One advantage of this “reality-in-a-test-tube” approach is that different intervention strategies can be tested beforehand at low cost with different model parameters for e.g. disease prevalence or contact tracing rate. Kretzschmar et al (1996) compared the effectiveness of different prevention and intervention scenarios for gonorrhea and Chlamydia, including contact tracing, mass screening, screening of subgroups and condom use.

In their simulations, they found that treatment of symptomatically infected and yearly screening of 20% of women in age class 15-24 was most effective in reducing Chlamydia prevalence. Treatment of at least 50% of partners was necessary to reduce Chlamydia prevalence to a low level with good probability of extinction (Kretzschmar et al, 1996). This shows how important contact tracing is for a long-term extermination program.

It should not be ignored, however, that a lot of the data involved in building models, choosing parameters and estimating starting values for simulations comes from different studies which are only related to each other via ecological correlation. Nevertheless, epidemiological models can help to highlight limitations in available information and to focus attention on what needs to be measured to better understand the complexity of infectious diseases (Garnett et al, 1996).

Literature was reviewed using Medline (2000) and Web of Science (2000) up to July 2000 and hand searching the journal Sexually Transmitted Infections up to September 2000. In addition, references were given by Pamela Warner, Dr. Sheena Sutherland and Dr. John Young. Further references were then taken from each article read. Keywords for online search were as follows:

Chlamydia specific:
Chlamydia trachomatis, Chlamydia, recurrence, recurrent infections, infection, reinfection.

Modelling:
stochastic/ deterministic/ theoretical model, sexually transmitted disease, simulation, Monte Carlo, computer, network, Markov Chain, modelling.
III. Study Design

Introduction

This dissertation seeks to answer important issues surrounding *Chlamydia* reinfections in patients of GUM clinics. More specifically, based on descriptive analysis and regression methods, the probability of reinfection within a time interval is estimated based on personal characteristics and testing history. This predicted reinfection interval would help clinicians to make the right recommendations for their patients and would also assist economists in making cost-benefit calculations for health service expenditures.

Study Population

The study cohort consists of all patients attending the Lothian GUM clinic between 1992 and May 2000. The Lothian Health Board is responsible for the health care needs of about 773,800 people living in the areas of East Lothian (89,600), Midlothian (80,900), West Lothian (153,100) and City of Edinburgh (450,200) (GROSa, 2000). They make up about 15% of the total Scottish population of 5,120,000 people (GROSa, 2000).

Data for this dissertation were derived from a review of *Chlamydia* test records between January 1992 and May 2000 for all patients attending the Royal Infirmary of Edinburgh GUM clinic (RIEGUM), the only one in Lothian. The clinic sees 9500 patients a year as of 1999-2000 (Dr. Gordon Scott, personal communication). The test records are stored in the Medical Microbiology Laboratory (MML) database, which is kept separate from the patients’ records database held at RIEGUM to ensure patients’ anonymity. Both databases can be record-linked. Data for this dissertation come from the MML database only, not the RIEGUM database.

The Royal Infirmary also serves as an outreach clinic for prostitutes (HEBS, 1999). In addition, its laboratories provide STD testing services for general practitioners (GPs) and family planning clinics. The ratio of number of tests done between GUM and non-GUM settings was 7376:4588 between 1.5.1999 and 1.4.2000. Non-GUM patients were almost exclusively women (Dr. Sheena Sutherland, affil.). Among the 7376 GUM patients, 3528 (48%) were women. However, this dissertation’s data exclude any tests from GPs or other non-GUM-settings.

Analysis of reinfection is further restricted to the subgroup of patients whose first positive test (=index test) was between January 1992 and December 1997 to ensure that everyone had at least 2.5 years time during which reinfections could be ascertained.
Routine Testing and Treatment Procedure

Patients coming to the RIEGUM clinic are given a unique patient identifier (UPI) at their first visit with the intention that this will be used for all future visits. The RIEGUM database stores information on name, date of birth (DOB), diagnosis, sex, ethnic group, postcode, reason for referral, occupational class, marital status, contraceptive method used and number of regular/irregular partners. Completeness of this data depends on a patient's cooperation and the comprehensiveness of a GP's referral. Unfortunately, the RIEGUM data was not available for this study.

Usually, patients are offered tests for Chlamydia and gonorrhea, even if they came for testing a different STD such as HIV. However, more Chlamydia than gonorrhea tests are made because of the convenience of giving a urine sample for Chlamydia compared to invasive probing for gonorrhea (Dr. Gordon Scott, personal communication). Specimens are then sent to the nearby MML where they get tested, usually on the same day. Each laboratory test gets a unique laboratory identifier (ULI). Test results are crosschecked by a senior scientific officer before being reported back to RIEGUM (Bruce Harris, personal communication). The MML uses the same patient identifier as RIEGUM, which enables record-linkage between both databases. Between January 1992 and August 1998 the large majority (98%) of Chlamydia tests was done by growth of culture and was superceeded from September 1998 onwards by ligase chain reaction (LCx) from Abbot Pharmaceuticals.

Patients are asked to come back after three days for the test results. In case of a positive test, they are additionally contacted by phone and asked to come back for treatment. If treated, they are further invited to return for a test of cure (TOC). The TOC should be made no sooner than four weeks after treatment. The reasons for this are twofold. First, antibiotics need enough time to kill pathogens and an early TOC could detect a bacterial population on the verge of eradication. Second, the new LCx tests are based on detecting DNA and minute amounts of undegraded DNA from dead bacteria could give a false positive result. This is likely to happen during the 2-3 weeks immediately after treatment.

Treatment follows the National Guideline for Chlamydia management (CEG, 1999) and consists of either 100mg Doxycycline twice a day for 7 days or a single 1000 mg dose of Acithromycine.

Reinfection Definition

same serovar, which could have been the result of either an incomplete cure (relapse) or an untreated partner.

Here, reinfection is defined as the second positive test of a patient and “reinfection interval” describes the time between the first and second positive test (fig. 2). An arbitrary number of negative tests may lie in between. The reinfection interval cannot be estimated exactly, because the first and second infection are likely to have occurred sometime before the tests by which each was detected.

In order to increase the probability of detecting new rather than uncleared previous (unresolved) infections, the time between two successive positive tests had to be equal or greater than 30 days. In case a patient tested positive within this 30-day interval, treatment failure was assumed and the next subsequent positive test, if done, was chosen.

Unresolved rather than true reinfections would be detected by a TOC about 2 months after treatment. However, this largely depends on a healthy patient's cooperation and of all 7,766 patients with two or more tests (total number of tests: 20,600), only 2,106 tests were carried out within 2 months. In order to make analysis easier, the stringent reinfection requirement of TOC used in prospective studies (Blythe et al, 1992, Fortenberry et al, 1999, Kjaer et al, 2000) was relaxed and both, patients' compliance and successful antibiotic treatment of first infection was assumed. 26 out of 34,754 patients had more than one reinfection episode, i.e. 3 or more positive tests. Multiple reinfection events will be discussed later.

Covariates in a Study

Covariates are used in regression modelling as independent factors to explain variations in outcome. The number of covariates that can be used in a regression calculation depends on the number of cases.
available and a range of other factors. The event per variable (EPV) ratio should be higher if there are small expected effects and dose-response gradients or if intercorrelations between variables or appreciable measurement errors exist.

Intraclass correlations, effect modification and heterogeneity of effects can further complicate modelling and may increase sample size needed (Camus, 2000). Unfortunately, a lot of these factors are not known until after data collection. If the EPV ratio is too small, the algebraic model that is used in proportional hazards regression might be unreliable and lead to spurious results (Concato et al, 1997), so inclusion of too many covariates should be avoided.

One difficulty for this study lies in the nature of laboratory data: it hardly contains any behavioural information on the patients other than GUM clinic visit patterns and the only physiological information stored are sex and age. Although information on ethnic group, postcode, occupational class, marital status, contraception used, number of regular and irregular partners is stored in the separate RIEGUM database, that information was not available at the time of this study.

Here, covariates based on visit pattern and test outcome were extracted and are used in addition to sex, age- and risk group. If too many variables are derived by indirect observations, they are likely to be strongly correlated with each other. This poses a methodological problem for any regression and choice of indirect covariates has to be carefully balanced. Details of covariates used are given in the next chapter.

The original study design consisted of an initial retrospective cohort study, with the intention of following it with a modelling simulation. The cohort study seeks to estimate the median reinfection interval for patients of GUM clinics and to assess the contribution of sex, age, risk group membership and GUM clinic visit history to reinfection risk. In addition, primary reinfection intervals will be compared with subsequent ones to see if they are different. Further, a comparison of diagnostic test performances tries to find out whether the new LCx test method has an influence on the likelihood of a positive test outcome. New DNA-amplification based tests are expected to pick up cases of infections that would have (false) negatives under the older methods (Young et al, 1998). This has not been proven yet on a large population level.

Finally, the modelling simulation would have to be built on the descriptive information of the data and would evaluate the performance of different contact tracing strategies. Modelling the efficiency of contact tracing strategies would have been done with Markov Chain Monte Carlo (MCMC) methods as simulation algorithms using WinBugs (Gilks, et al, 1996, BUC/DEPICL, 2000). A fundamental assumption is that the probability of an event is independent of event history (Markov property), i.e. the probability for an individual testing positive for Chlamydia does not depend on the outcomes of previous Chlamydia tests. In the event the cohort study only was available in the timeframe for the dissertation, the simulations will be conducted at a later date.
IV. Methods

Estimation of Reinfection Intervals

Survival methods are used to estimate the time to reinfection and factors contributing to risk of reinfection. They allow for incomplete observations and different starting points for observations through time and record the time interval between start of observation and the event happening. In this context, the statistical term “event” represents reinfection with *Chlamydia* and the term “survival” corresponds to the time to event, not a patient’s actual survival.

“Censoring” of observation happens in patients who either withdrew from study without having had the event or who have had no event during the whole study period. Right censoring relates to withdrawal from study, left censoring occurs when the starting point of a person's time-to-event is not precisely known (Kleinbaum, 1995). This is the case for most survival studies involving infections: the exact time of infection is never known, only the time of the first positive test.

Survival methods assume that entering or being withdrawn from follow-up in a study is unrelated to the current hazard of the event happening (non-informative censoring), otherwise systematic inclusion or withdrawal of high- or low-risk patients would bias the results (Bull *et al* 1997). However, if the event occurred in a high-risk patient at the beginning of the study in 1992-93 and that person then had all subsequent tests at a GP instead of the GUM clinic, the withdrawal was related to the reinfection hazard. This situation cannot be controlled for with GUM clinic data only. Systematic withdrawal is termed “informative right censoring”. “Informative left censoring” can happen if late arrivals into the study are not at equal risk to those already under surveillance (non-informative late entry) (Bull *et al* 1997).

Because this study is based on routine data, there is no reason to believe otherwise.

Kaplan-Meier (KM) curves (survival curves) plot against time the probability that a study subject survives, i.e. is event-free past a specified time (Kleinbaum, 1995). They are graphical representations of life tables, which record the time between events and the proportion of event-free patients. The median time-to-event, here median reinfection time, can be obtained graphically by looking at which time the survival-probability equals 0.5, i.e. passes through y=0.5. This estimation is only reliable if the survival curves falls rather steeply through y=0.5. KM curves can be plotted for different levels of a factor, e.g. sex, and equality of survival distributions for the different levels can be tested with the logrank test (Bull *et al*, 1997). It requires constant odds ratios of risk through time, i.e. constant slopes of survival curves. In this study, the distributions between men and women, stratified for age, is compared.
To test the influence of more than one covariate on time-to-event, more complex methods have to be chosen. Cox's proportional hazards regression model will be used to evaluate risk factors for reinfection. A regression model looks at adjusted influences of specific factors on outcome and tries to predict (within limits) the outcome for an individual with a certain set of characteristics. Cox’s proportional hazards regression model is a technique that provides simultaneous estimates of hazard ratios in the presence of multiple explanatory factors (Bull et al 1997). It is a semiparametric model and expresses the instantaneous risk of an event occurring (=hazard) as a parametric function of the factors of interest (covariates) multiplied with an underlying non-parametric baseline hazard function for the event. Time to event models that permit analysis of multiple events per subjects (multistate hazard models), i.e. patients with more than one reinfection, are currently topics of discussion in statistical research (Clayton, 1994, Gordon Murray, personal communication) and will not be used here. In addition, standard statistical software packages such as SPSS do not support them, yet. Therefore, only the first reinfection-event will be used for survival analysis.

Cox’s regression model assumes that covariates (e.g. sex or age) have a multiplicative effect on the hazard function and that the ratio of hazard functions for any two individuals will be constant through time, i.e. the covariates included in the model are independent of time (proportional hazards). Cox’s model does not make any assumptions about the underlying hazard functions other than being proportional. To test the assumption of proportional hazards, one could build a more complex model with a time dependent covariate and look whether the time dependent factor is significant or not. One could also divide the time into different epochs, make a Cox’s regression for each single epoch and then check whether the covariates’ coefficients differ markedly.

Fortunately, one can check the assumption of proportional hazards graphically with a “log(-log of survival function)”-plot (LML-plot) against time for a number of subgroups defined by different combination of covariates, which is the method of choice in this dissertation. If the assumption holds, the plots should produce a number of parallel lines. First, a univariate LML plot has to be made for different values of each single covariate involved in the selection process. This is followed by LML plots for all different combinations of significant covariates. In case a covariate violates the assumption of proportional hazards, the analysis can be split up into subanalyses, stratified for this covariate. It is also informative at what period in time the assumption was violated.

Cox’s model also allows assessing the impact on time to an event of a particular covariate, adjusted for the other covariates. For example, the influence of a patient's sex on reinfection risk can be assessed independent of age.
Population and Covariates included in Cox’s Regression

To enter the study, patients had to have a positive test between 1992 and 1997 and a subsequent negative or positive test (n=1610). For reinfecteds, time to event (=reinfection) was calculated as: (“date of second positive test” – “date of index test”), non-reinfected patients had their time to censoring calculated as: (“date of last negative test” – “date of index test”). The subsequent positive test had to be more than one month apart, which was not the case for 21 patients. 3 of these 21 patients had no more tests done and were excluded. Another 3 of the 21 cases had a third positive test one or more months after the index test, which was then taken to calculate the reinfection time. The remaining 15 patients were treated as having had one positive and one or more negative tests, i.e. as not reinfected. This made a total of n=1607 cases which corresponds to group D2 in figure 3 (appendix). The age distribution per sex of the 1607 patients will be compared by a WRS-test with the remaining study population to see whether they are similar and results can be extrapolated.

Agegroup and sex are extracted directly from the data. Agegroups are defined according to the agebands used by ISD (ISD, 2000). Agegroup is used as a nominal instead of age as a continuous variable because it is known that for men, *Chlamydia* incidence first rises with age and then decreases. This clearly violates the assumption of linear effects on on hazard for a continuous variable. However, one must realize that using several nominal variables instead of one continuous reduces the statistical power of a regression, so their number should be minimized. The following agegroups are chosen: 15-19, 20-24, 25-34, ≥35. The first three correspond to those chosen by ISD (2000), the fourth summarises the last two ISD-agebands.

Patients from the outreach clinic in Leith are prostitutes and have a certain letter in their UPI. As they comprise a defined risk group, a binary variable called prost will be included in the analysis and set to one if the patient is a prostitute. It is not clear, however, whether they are more at risk of acquiring a STD since they can counteract this “occupational” risk by insisting on condom-use.

Trends of reinfection through time could be measured by including year of index case as a covariate. If taken as a continuous variable, one assumes a linear effect on reinfection risk. It seems more likely, however, that risk behaviour changed abruptly because of HIV and Safe Sex campaigns. Unfortunately, there were no major campaigns in Lothian during the study period (Dr. Gordon Scott, personal communication). Accounting for year of test without making invalid assumptions would require 6 additional nominal variables (1992-1997), at considerable cost to the statistical power of the study. It is therefore not included. Year of test still is an important covariate in a study, especially if a shift in tests towards amplification assays occurred during the study period.
Estimating Reinfection Intervals for *Chlamydia trachomatis*

One could also take year of first visit at the clinic as a behavioural covariate. It is not used here either for the same reasons given for year of test.

People who come to a GUM clinic for the first time come for a reason, usually they had risked exposure or have symptoms. In case of a symptomatic patient, the person will likely test positive on the first visit and might be more careful in the future, thus increasing the interval to reinfection. On the other hand, someone who tests negative on the first visit might get a complacent attitude towards sexual risk behaviour and have a shorter reinfection interval. To test this, a binary variable will be included in the regression (ta. 2). It is set to 1 if a reinfected patient had one or more negative tests prior to the index test. Total number of clinic visits is not included as a covariate because in 66% of the cases with 3 or more visits the additional visit(s) took place after the index case and would thus not be known beforehand, which makes “number of visits” less suitable as a prognostic factor. It would also be strongly correlated with a variable measuring prior negative tests, as the number of total visits rises with the likelihood of a previous negative visit.

Summing up, covariates used in Cox’s regression are age category, sex, risk group membership and visit history (tab. 2). Variable names used in the text will appear in “Courier” font.

Table 2: Covariates used in Cox’s regression.

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Comparison of patients with one vs. multiple clinic visits

Routine GUM data depends on patients coming for testing voluntarily and a reinfection can only get picked up if they have at least 2 visits. The group of patients with one visit only could be systematically
different from the group with two or more visits, which would bias any results regarding reinfection intervals.

To detect differences, the age at first visit of both sub-populations will be compared for each sex with a Wilcoxon Rank Sum (WRS)-test, the null-hypothesis being that of no difference between the groups. The non-parametric WRS test is chosen since the distribution of age per sex and patient group is not known. Only patients who presented before 1998 will be chosen to allow everyone at least 2.5 years time to return to the clinic.

Comparison of Multiple Reinfection Episodes

Cox's regression model described above was developed for non-repetitive events such as death. Multiple events such as successive reinfections can not be included in basic Cox's regression analysis. In order not to discard potentially useful information on reinfection, the length of subsequent reinfection intervals will be compared with that of primary reinfection intervals.

A person with a Chlamydia reinfection might have become more responsible in his or her sexual risk behaviour and have longer subsequent reinfection intervals. Alternatively, since Chlamydia is easily cured with antibiotics, a person's perception of STDs might be that of a minor nuisance, conquered by modern technology and intervals would shorten. Insight into these patterns could help targeting education and screening efforts. Patients will be older at the second interval by definition, and should age be associated with a decreased risk of reinfection, any secondary reinfection interval would thus tend to be longer. Comparing the intervals of patients with multiple reinfections could pick up differences in length of intervals between first and subsequent reinfections. Due to the low numbers of patients with more than one reinfection episode (26 out of 34,754 patients), only first and second reinfection intervals will be compared. Distribution of reinfection intervals is unknown, so a Wilcoxon signed rank test for paired samples will be used. The null hypothesis is that intervals of secondary reinfections do not differ from those of the first. The age distribution per sex of the multiple-reinfection subgroup then has to be compared by a WRS-test with the general study population to check generalisability.

Impact of increased test sensitivity

Given a constant risk of infection, incidence rates would go up automatically if a more sensitive test is used and by only looking at the rates one would assume an increase of risk. With regard to the MML Chlamydia data from 1992 to 2000, on September 1998 a switch in testing methods from culture to LCx occurred. DNA amplification assays have a higher sensitivity compared to culture methods (Young et al,
1999) and thus incidence and reinfection rate would be expected to rise after September 1998 because of the new test only. To check this on a large population scale, proportion of positive diagnoses for women undergoing cervical smear tests will be compared by Chi-square-test one year before (group 1) and after (group 2) the change in tests. The nullhypothesis is that both proportions are equal. Again, age will be compared between both groups by a WRS-test to test their homogeneity.

Statistical Analysis

The estimation of reinfection intervals, the tests for comparison of patients with one vs. multiple clinic visits, comparison of multiple reinfection episodes and increased test sensitivity will be made as described above. They will be preceded by a descriptive analysis of the MML data. First, general population characteristics of the test-based MML data will be given with respect to sex and age of patients and compared to the composition of the general population in Lothian. Then, a pivot table will describe the number of *Chlamydia* tests and their outcomes for each year, stratified by sex and ageband. It is followed by graphs of the number of positive and negative tests per ageband, stratified by sex and year of test. The graphs do not contain additional information, however, they help to better visualise *Chlamydia* incidence per sex and ageband through time.

The sub population used in the survival analyses will then be described in more detail by giving the proportion of patients reinfected within one, two and three years, stratified for sex and ageband. Finally, a plot of the proportion of positives against age at testing for men and women will illustrate age trends in infection between the sexes.
V. Data management

Data Cleaning

Data cleaning is an essential and often overlooked issue of epidemiological research. It describes the techniques necessary to resolve inconsistencies within the dataset. In the case of retrospective studies such as this one, data often come from routinely collected information over many years and one has usually little control over the collection process. It cannot automatically be assumed that the dataset is free of errors and inconsistencies. Systematic differences during the data collection may lead to recording bias. Further, any loss of quality of the data weakens the statistical inferences drawn and might mask true associations or create spurious ones between the variables of interest. Diagnostic errors are exceptionally difficult to detect afterwards and only strict laboratory quality control can prevent them from happening in the first place. Errors made during electronic data entry, e.g. regarding sex or DOB can show up as inconsistencies if the false information on one record can later be matched through a database with that from a correct one. Some errors happen because of poor design of report sheets or user interfaces. With the growing number of retrospective studies based on electronic archives of patients' records and multi-centre databases, data cleaning techniques will become more and more important.

Data storage system

From 1992 on, test results on all STD tests were kept electronically in a database. These records contain the UPI, DOB, sex, location of specimen, date of sampling, date of testing, ULI, setting (RIEGUM, GP) and comments with test results. Information on STDs other than Chlamydia and reason for visit for some patients (e.g. termination of pregnancy, cervical cancer screening) were stored in the MML database but were not extracted for this analysis.

Information regarding a patient's place of residence, ethnic group, occupational class, number of regular/irregular sex partners, contraception used and marital status is not stored but could be retrieved by record linkage from the RIEGUM database.

Multiple MML records can be crosslinked via UPIs to create summary reports, e.g. on all Chlamydial and gonococcal tests an individual has had (Bruce Harris, personal communication).
Data extraction and cleaning

MML test records are stored in two different databases, one for tests done before summer 1998 (40716 records) and one for tests done thereafter (8175 records). Data for this study have been extracted from both systems into two Microsoft Access files. The database system used in this dissertation was FilemakerPro 4.0 for Macintosh. Both files were transferred from Access to FilemakerPro via DBF 4.0 format, which is supported by both programs. Data transfer consistency has been checked by comparing total tests done, total number of females and total number of males.

Minor adjustments had to be made to the original data. The old MML database system allowed for multiple comments per entry and each extra line of comment created an additional record if exported to Access. This led to several entries (1.273 of 40.716) with the same laboratory identifier, violating its uniqueness. After manual elimination of records with duplicate ULIs, both data files were concatenated and resulted in 47.618 entries (40.716+8.175-1.273). Result and type of test were extracted from the comment field into new variables. Age at test was calculated by subtracting DOB from date of test. For a table of variables in the database, refer to table 3 (appendix).

In summer 1998, MML switched to a new database system

One problem were records with the same patient identifier, but different DOB (351) and/or sex (53), i.e. tests seeming to be of the same patient, but discordant as to DOB or sex (400 of 47.618/ 0.8%). Some DOB-differences (124) were of typographical nature, where e.g. a “3” in one record became an “8” in another or a “1” became a “7”. Only one digit in either day, month or year was discordant. Other records (227) had completely different DOBs with different day, month and year. Here, either DOB was correct and the UPI was incorrectly read from the request form (appendix, fig. 4.1), so a genuinely different person was tested. Alternatively, a flaw in the user interface could have lead to the same person getting a wrong DOB: usually, UPI, DOB and sex have to be entered by a lab technician. If only UPI is entered, DOB and sex of the previously entered patient remains on the screen and is, erroneously, used. A higher proportion of these non-typographical errors occurred after the database switch in summer 1998 (appendix, fig. 5), maybe because records prior to the switch were not accessible from the new database and thus proofreading was not possible. Errors also doubled from the third quarter 1997 onwards, maybe due to a change in data entry staff. However, the more tests per UPI are made and the larger a database is, the more likely these errors can arise.

Typographical DOB conflicts were resolved “democratically” with the DOB of the majority of records with the same UPI overruling the discordant record. If there were only two records with the same UPI
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(95), DOB of that patient was looked up at the original MML database, which included patient entries other than that for *Chlamydia*. If there were no further entries, the later test was not used.

For records with same UPIs but completely different DOBs, those in the minority were excluded. In case of a draw, the later presentation was excluded. A total of 227 tests have been excluded from analysis this way. This might have introduced bias and will be discussed later.

Records with conflicting sex (53 of 400) were treated differently. The request sheet that accompanies each specimen when it arrives at the lab has two fields for sex, male and female which lie close together and by circling the field hastily or carelessly, the wrong field can easily be indicated. Fortunately, UPIs consist of a letter and a four-digit number. 9 of 11 letters code for either male or female, so all except 12 sex-conflicts could be resolved with the UPI-keys (Bruce Harris, personal communication).

Some records had internal inconsistencies such as men with cervical swabs. However, the location-field on the request form had the boxes for “urethra” and “endocervix” next to each other (appendix, fig 4.2), so a hastily filled out form might have resulted in the wrong box indicated. These inconsistencies were ignored, because “location of specimen” was not used in the analysis. Records with *Chlamydia* tests for eyes (n=29) were excluded from analysis, since they were unlikely to be transmitted sexually. Altogether, 256 out of 47618 records (0.5%) were excluded.

It is important to remember that each record comprised one test, so the original per-test database had to be converted into a per-patient database for further analysis. In the latter, each record corresponds to one patient (UPI) with additional information such as total number of tests, test outcomes, number of positive and negative results or interval between first positive test and second positive test. Extracts from the total dataset were made for the individual statistical analysis.
VI. Results

Descriptive Analysis

The age distribution of patients visiting the RIEGUM clinic differs for men and women (fig. 6.1). In both sexes, there were very few patients under 15 years of age, but apart from this, female patients were much more likely to be under 25 years (53% of women patients compared to 32% of men). Men dominated the 25+ agebands.

![Age distribution for men and women](image)

Figure 6.1: Number of total Chlamydia tests for men (blue) and women (red) for each age category.

Figure 6.2 compares the relative proportion of men and women in different agebands between the RIEGUM clinic population and the general Lothian population (GROSa, 2000). People between 15-29 dominate the sexually active population in Lothian, they are about three times more abundant in the RIEGUM data set than in the general population.
Proportion of men and women per ageband compared between Lothian and RIEGUM population

Figure 6.2: Comparison of relative age distribution between the RIEGUM clinic population and the general Lothian population.

The pivot table lists the number of men testing positive or negative and the number of women testing positive or negative for each year between January 1992 and May 2000 (appendix, tab. 4). The numbers are further given separately for the different age-categories. The categories were set according to those used in ISD publications, however with categories 45-64 and ≥65 put together (ISD, 2000). A total of 47589 *Chlamydia* tests were done in that period. Only 47305 records are given in the pivot table because few tests had no sex (12) and/or no DOB (51) or were marked having unreliable results in the MML database (223).

In order to get a better overview, the “outcome per ageband” information is summarised for the years 1992 - 1999 in figure 7 on a semi-log scale, where the average number of positive and negative test outcomes in men and women including standard deviation is given for each ageband. Note that number of tests is plotted on a log-scale so that agebands with low numbers (0-14) and high numbers (25-34) can be displayed on the same graph.

Standard deviation is chosen as dispersion measure because the sample comprises the entire study population. It can be seen that negative tests always outweigh positive tests in both sexes and all agebands. The number of positive cases for women is greater than that in men in the 15-19 ageband, about the same in the 20-24 ageband and consistently lower in the upper agebands.
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**Figure 7**: Tests per ageband and sex. Summary of the information of table 4 (pivot table). The 8 years 1992-1999 were combined to give the average number of positive and negative *Chlamydia* tests for men (dark and light blue) and women (dark and light red) per ageband. Graph includes standard deviation (black error bars). Number of tests is given on a logarithmic scale.

Figures 8.1-8.4 (appendix) contain the same information given in the pivot table, but in a different arrangement. Here, the number of men and women testing positive or negative are displayed per ageband on a timescale from 1992 to 1999 to look at trends in infections during that period. Although it is a bit difficult to compare graphs on a semi-log scale, it can be seen clearly that women have consistently more positive cases than men in the 15-19 ageband and less in the 25-34 and 35-44 ageband. With regard to test outcomes in the 20-24 ageband, men and women are virtually indistinguishable. It can further be seen in the graphs that the number of men and women testing positive increased from 1997 onwards, but so did the number of men and women testing negative.

Table 5 describes the population used in the survival analysis (group D2 in appendix, fig. 3) with regard to reinfection within one, two, three and four or more years. The total number and proportion of reinfected men and women is given four age-categories. The categories used here differ slightly from the six used above, in that the first (0-14) is omitted and the last two (35-44, ≥45) are combined.
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Group men: 15-19 and group women: ≥35 contain less than 30 cases, so their results have to be treated with caution. In the other agegroups, proportion of reinfected within one year is between 2.0% and 4.8% for men and 3.2% and 8.8% for women. The women: 15-19 group has the highest proportion (8.8%) of reinfected within one year. Within three years, 7.7% of men between the age of 20-24 and 12.5% of women between 15-19 become reinfected with *Chlamydia*.

Table 5: Reinfection status after one, two, three and four or more years per sex and agegroup for the population used in the survival analyses.

<table>
<thead>
<tr>
<th>sex, age</th>
<th>patients</th>
<th>Reinfected within</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>one year</td>
<td>two years</td>
<td>three years</td>
<td>four or more years</td>
</tr>
<tr>
<td>Men</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>15-19</td>
<td>27</td>
<td>1 (3.7%)</td>
<td>1 (3.7%)</td>
<td>2 (7.4%)</td>
<td>3 (11.1%)</td>
</tr>
<tr>
<td>20-24</td>
<td>309</td>
<td>15 (4.8%)</td>
<td>20 (6.4%)</td>
<td>24 (7.7%)</td>
<td>30 (9.7%)</td>
</tr>
<tr>
<td>25-34</td>
<td>445</td>
<td>18 (4.0%)</td>
<td>24 (5.3%)</td>
<td>26 (5.8%)</td>
<td>37 (8.3%)</td>
</tr>
<tr>
<td>35-100</td>
<td>97</td>
<td>2 (2.0%)</td>
<td>4 (4.1%)</td>
<td>4 (4.1%)</td>
<td>5 (5.1%)</td>
</tr>
<tr>
<td>total</td>
<td>878</td>
<td>36 (4.1%)</td>
<td>49 (5.6%)</td>
<td>56 (6.4%)</td>
<td>75 (8.5%)</td>
</tr>
<tr>
<td>Women</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>15-19</td>
<td>136</td>
<td>12 (8.8%)</td>
<td>15 (11.0%)</td>
<td>17 (12.5%)</td>
<td>19 (14.0%)</td>
</tr>
<tr>
<td>20-24</td>
<td>358</td>
<td>14 (3.9%)</td>
<td>15 (4.1%)</td>
<td>15 (4.2%)</td>
<td>18 (5.0%)</td>
</tr>
<tr>
<td>25-34</td>
<td>214</td>
<td>7 (3.2%)</td>
<td>8 (3.7%)</td>
<td>9 (4.2%)</td>
<td>11 (5.1%)</td>
</tr>
<tr>
<td>35-100</td>
<td>21</td>
<td>1 (4.7%)</td>
<td>1 (4.7%)</td>
<td>1 (4.7%)</td>
<td>1 (4.8%)</td>
</tr>
<tr>
<td>total</td>
<td>729</td>
<td>34 (4.6%)</td>
<td>39 (5.3%)</td>
<td>42 (5.8%)</td>
<td>49 (6.7%)</td>
</tr>
</tbody>
</table>

In figure 9, the proportion of women and men testing positive is plotted against age at infection to look at trends between the sexes. Proportion of positives under 16 years is unreliable because of the low total number of tests done. For both sexes, there is an overall decrease from 15% down to 0-3%. 17-year-old women (14.7%) have four percent points more positive tests than men (10.6%) of that age. Both sexes are roughly equal with 18, and from 22 years on men have consistently 2-4 percent points more positive tests than women.
Hypothesis Testing

Survival Analysis

The age distribution of men and women in the sub-population used for the survival analyses was significantly different from that of the general GUM population (p<0.0001 for men and women). Median age of men was 25.3 compared to 28.2 in the general population, 21.6 years compared to 24.8 for women (tab. 6).

Table 6: Results of testing the null hypothesis that the age distribution is the same for men and women in the survival-study population compared to the general population.

<table>
<thead>
<tr>
<th></th>
<th>survival-study pop.</th>
<th>remaining population</th>
<th>Z value of 2 tailed WRS-test</th>
<th>2-tailed significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>number of men</td>
<td>878</td>
<td>10956</td>
<td>-11.6</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>median age</td>
<td>25.3</td>
<td>28.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>number of women</td>
<td>729</td>
<td>9866</td>
<td>-14.3</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>median age</td>
<td>21.6</td>
<td>24.8</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
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The Kaplan-Meier plot for men and women of all ages (fig. 10.1) shows that in the early months women are more at risk of reinfection, however, after about 18 months the instantaneous risk of reinfection is higher for men than for women. Separate plots for the agebands 15-19, 20-24, 25-34 and ≥35 years give a similar picture (fig. 10.2-5), but there are too few men in ageband 15-19 and too few women in ageband ≥35 to give reliable plots. Median times to reinfection in months for men were 67 (overall age), 64 (agecat:15-19), 60 (agecat:20-24) and 80 (agecat:25-34). For women, it was 54 (agecat:15-19) and 77 months (agecat:25-34). The survival curves for women:20-24, women:≥35, women:all and men:≥35 never went below y=0.5 and thus gave no median survival time.

![Survival Functions, all years](image)

Figure 10.1: Kaplan-Meier plot of cumulative survival for men and women of all ages. Y-axis begins with 0.3.
Estimating Reinfection Intervals for *Chlamydia trachomatis*

**Survival Functions, 15-19 years**

![Kaplan-Meier plot](image)

**Survival Functions, 20-24 years**

![Kaplan-Meier plot](image)

Figure 10.2: Kaplan-Meier plot of cumulative survival for men and women between 15 and 19 years.

Figure 10.3: Kaplan-Meier plot of cumulative survival for men and women between 20 and 24 years.
Figure 10.4: Kaplan-Meier plot of cumulative survival for men and women between 25 and 34 years, y-axis begins with 0.2.

Figure 10.5: Kaplan-Meier plot of cumulative survival for men and women 35 years and older, y-axis begins with 0.4.
The logrank test was used to check equality of survival distribution between men and women in the different agebands (tab. 7). Only in the 20-24 ageband a significant difference was detected.

Table 7: Logrank tests to test equality of survival distributions between man and women. Tests are made for all ages and each individual age-category.

<table>
<thead>
<tr>
<th>age category</th>
<th>sex</th>
<th>median time to reinfection (months)</th>
<th>number of reinfections</th>
<th>number censored</th>
<th>Log Rank</th>
<th>significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>15-19</td>
<td>men</td>
<td>64</td>
<td>3</td>
<td>24</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>women</td>
<td>54</td>
<td>19</td>
<td>117</td>
<td>0.32</td>
<td>0.5720</td>
</tr>
<tr>
<td>20-24</td>
<td>men</td>
<td>60</td>
<td>30</td>
<td>279</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>women</td>
<td>#</td>
<td>18</td>
<td>340</td>
<td>4.66</td>
<td>0.0309</td>
</tr>
<tr>
<td>25-34</td>
<td>men</td>
<td>80</td>
<td>37</td>
<td>408</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>women</td>
<td>77</td>
<td>11</td>
<td>203</td>
<td>1.09</td>
<td>0.2974</td>
</tr>
<tr>
<td>35-100</td>
<td>men</td>
<td>#</td>
<td>5</td>
<td>92</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>women</td>
<td>#</td>
<td>1</td>
<td>20</td>
<td>0.00</td>
<td>0.9867</td>
</tr>
<tr>
<td>all</td>
<td>men</td>
<td>67</td>
<td>75</td>
<td>803</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>women</td>
<td>#</td>
<td>49</td>
<td>680</td>
<td>1.07</td>
<td>0.3002</td>
</tr>
</tbody>
</table>

# survival curve (survival probability) did not fall below 0.5

Frequencies for covariates used in the Cox’s regression are given in table 8. 10% of all study subjects are between 15 and 19 years old, 41% between 20-24 and 25-34, 8% are ≥35. 12% had a negative test before the index test and 0.7% are prostitutes.
Table 8: Covariates used in Cox’s regression.

<table>
<thead>
<tr>
<th>covariates used in Cox’s regression</th>
<th>name in regression</th>
</tr>
</thead>
<tbody>
<tr>
<td>age category</td>
<td>name in regression</td>
</tr>
<tr>
<td>15-19</td>
<td>agecat15-19</td>
</tr>
<tr>
<td>20-24</td>
<td>agecat20-24</td>
</tr>
<tr>
<td>25-34</td>
<td>agecat25-34</td>
</tr>
<tr>
<td>≥35</td>
<td>agecat≥35</td>
</tr>
<tr>
<td>1st test negative</td>
<td>firstneg</td>
</tr>
<tr>
<td>yes</td>
<td>1</td>
</tr>
<tr>
<td>no</td>
<td>0</td>
</tr>
<tr>
<td>prostitute</td>
<td>prost</td>
</tr>
<tr>
<td>yes</td>
<td>1</td>
</tr>
<tr>
<td>no</td>
<td>0</td>
</tr>
</tbody>
</table>

The LML plots for sex (fig. 11.1) and firstneg (fig. 11.2) have non-parallel curves, the plot for prost has one curve only (fig 11.3). The LML plot for agebands has parallel lines except for agecat≥35 (fig. 11.4).

Figure 11.1: Log-minus-Log plots to check proportional hazards assumption for covariate sex.
Figure 11.2: Log-minus-Log plots to check proportional hazards assumption for covariate firstneg.

Figure 11.3: Log-minus-Log plots to check proportional hazards assumption for covariate prost.
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### Figure 11.4: Log-minus-Log plots to check proportional hazards assumption for covariate agecat.

Three Cox's regressions were made, one (Cox1, tab. 9.1) with the variables \textit{sex}, \textit{prost} (=prostitutes), \textit{agecat} and \textit{firstneg} (=negative test prior to index test) and two for each, men (Cox2, tab. 9.2) and women (Cox3, tab 9.3) with the variables \textit{prost}, \textit{agecat} and \textit{firstneg} only. The extra two regressions became necessary because \textit{sex} violated the proportional hazards assumption (see discussion).

Table 9.1: Cox’s regression with covariates \textit{sex}, \textit{agecat}, \textit{firstneg}, \textit{prost}. 760 cases of 1607 were excluded from regression modelling because they were censored before the first event happened.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coefficient</th>
<th>S.E.</th>
<th>Significance</th>
<th>Exp (b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>sex</td>
<td>-0.3768</td>
<td>0.2025</td>
<td>0.0628</td>
<td>0.686</td>
</tr>
<tr>
<td>pros</td>
<td>-10.8934</td>
<td>191.4236</td>
<td>0.9546</td>
<td>1.86E-05</td>
</tr>
<tr>
<td>agecat</td>
<td>0.0289</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>15-19</td>
<td>0.6201</td>
<td>0.2149</td>
<td>0.0039</td>
<td>1.8591</td>
</tr>
<tr>
<td>20-34</td>
<td>0.0472</td>
<td>0.1652</td>
<td>0.7751</td>
<td>1.0483</td>
</tr>
<tr>
<td>25-34</td>
<td>-0.164</td>
<td>0.1662</td>
<td>0.3237</td>
<td>0.8487</td>
</tr>
<tr>
<td>≥35</td>
<td>0.0205</td>
<td>0.234</td>
<td>0.9301</td>
<td>1.0207</td>
</tr>
</tbody>
</table>
Table 9.2: Cox’s regression for men only with covariates agecat, firstneg, prost. 451 cases of 878 were excluded from regression modelling because they were censored before the first event happened.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coefficient</th>
<th>S.E.</th>
<th>Significance</th>
<th>Exp (b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>prost</td>
<td>-6.5494</td>
<td>308.2868</td>
<td>0.9831</td>
<td>1.40E-03</td>
</tr>
<tr>
<td>agecat</td>
<td></td>
<td>0.6476</td>
<td></td>
<td></td>
</tr>
<tr>
<td>15-19</td>
<td>0.2091</td>
<td>0.4575</td>
<td>0.2429</td>
<td>1.2326</td>
</tr>
<tr>
<td>20-34</td>
<td>0.2746</td>
<td>0.2351</td>
<td>0.8069</td>
<td>1.316</td>
</tr>
<tr>
<td>25-34</td>
<td>-0.0552</td>
<td>0.2257</td>
<td>0.3237</td>
<td>0.9463</td>
</tr>
<tr>
<td>≥35</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>firstneg</td>
<td>0.022</td>
<td>0.3071</td>
<td>0.9429</td>
<td>1.0223</td>
</tr>
</tbody>
</table>

Table 9.3: Cox’s regression for women only with covariates agecat, firstneg, prost. 309 cases of 729 were excluded from regression modelling because they were censored before the first event happened.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Coefficient</th>
<th>S.E.</th>
<th>Significance</th>
<th>Exp (b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>prost</td>
<td>-11.9822</td>
<td>341.5366</td>
<td>0.972</td>
<td>6.25E-06</td>
</tr>
<tr>
<td>agecat</td>
<td></td>
<td>0.0476</td>
<td></td>
<td></td>
</tr>
<tr>
<td>15-19</td>
<td>0.6418</td>
<td>0.3253</td>
<td>0.0485</td>
<td>1.8998</td>
</tr>
<tr>
<td>20-34</td>
<td>-0.1666</td>
<td>0.3255</td>
<td>0.6087</td>
<td>0.8465</td>
</tr>
<tr>
<td>25-34</td>
<td>-0.2123</td>
<td>0.3508</td>
<td>0.5451</td>
<td>0.8087</td>
</tr>
<tr>
<td>≥35</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>firstneg</td>
<td>0.0098</td>
<td>0.3622</td>
<td>0.9783</td>
<td>1.0099</td>
</tr>
</tbody>
</table>
Estimating Reinfection Intervals for *Chlamydia trachomatis*

The risk of reinfection within a month’s time, assuming non-infection up to \( t \) would be:

**Cox1:**

\[
h(t) = h_0(t) \times \exp(-0.377 \times \text{sex} - 10.893 \times \text{prost} + 0.021 \times \text{firstneg} + 0.620 \times \text{agecat15-19} + 0.047 \times \text{agecat20-24} - 0.164 \times \text{agecat25-34})
\]

**Cox2:**

\[
h(t) = h_0(t) \times 0.686 \times \text{sex} \times 0.00002 \times \text{prost} \times 1.021 \times \text{firstneg} \times 1.859 \times \text{agecat15-19} \times 1.048 \times \text{agecat20-24} \times 0.849 \times \text{agecat25-34}
\]

**Cox3:**

\[
h(t) = h_0(t) \times \exp(-6.549 \times \text{prost} + 0.022 \times \text{firstneg} + 0.209 \times \text{agecat15-19} + 0.275 \times \text{agecat20-24} - 0.055 \times \text{agecat25-34})
\]

\[
h(t) = h_0(t) \times 0.001 \times \text{prost} \times 1.022 \times \text{firstneg} \times 1.233 \times \text{agecat15-19} \times 1.316 \times \text{agecat20-24} \times 0.946 \times \text{agecat25-34}
\]

**Cox3:**

\[
h(t) = h_0(t) \times \exp(-11.982 \times \text{prost} + 0.010 \times \text{firstneg} + 0.664 \times \text{agecat15-19} - 0.167 \times \text{agecat20-24} - 0.212 \times \text{agecat25-34})
\]

\[
h(t) = h_0(t) \times 0.000006 \times \text{prost} \times 1.010 \times \text{firstneg} \times 1.900 \times \text{agecat15-19} \times 0.846 \times \text{agecat20-24} \times 0.809 \times \text{agecat25-34}
\]

with coding according to table 8.

Cox1 had only \text{agecat} as a significant covariate (\( p=0.0289 \)), more specifically only ageband 15-19 was significant (0.0039). \text{sex} was borderline non-significant with \( p=0.0686 \). Cox2 had no significant covariates and in Cox3, only \text{agecat} was significant (\( p=0.0476 \)). Again, only ageband 15-19 had a \( p \)-value under 5% (\( p=0.0485 \)). SPSS reported that during calculation of the Cox1 regression model, 760 out of 1607 cases had to be dropped because censoring occurred before the earliest event in a stratum. Cox2 had 451 dropped out of 878, Cox3 309 out of 729. The implications of this will be discussed below.
Patients with one vs. multiple clinic visits

Men and women going to the clinic once differ statistically significant in age from those going twice or more (tab. 10).

Table 10: Results of testing the nullhypothesis that men and women with one visit have the same age-distribution as those with two or more visits.

<table>
<thead>
<tr>
<th></th>
<th>patients with 1 visit only</th>
<th>patients with 2+ visits</th>
<th>Z value of 2 tailed WRS-test</th>
<th>2-tailed significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of men</td>
<td>8666</td>
<td>3168</td>
<td>-4.58</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Median age</td>
<td>28.1</td>
<td>27.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of women</td>
<td>7687</td>
<td>2908</td>
<td>-12.8</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Median age</td>
<td>25.0</td>
<td>23.4</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Multiple Reinfecion Episodes

The data shown in table 11 are consistent with the nullhypothesis of no difference between first and second reinfection interval (p=0.919, WRS). However, the age distribution between the general GUM population and patients with multiple reinfection intervals is significantly different. Men and women with multiple reinfections tend to be younger than the general GUM population (tab. 4).

Table 11: Results of testing the nullhypothesis that the durations of first and second reinfection intervals are the same. Also given is the result of testing whether the age distribution of men and women in the preceding test is different from that of the other GUM clinic patients.

<table>
<thead>
<tr>
<th>Wilcoxon matched-pairs Signed Rank Test of first and second reinfection interval (n=21)</th>
<th>1st longer than 2nd</th>
<th>2nd longer than 1st</th>
<th>Z-score</th>
<th>2 tailed sign.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>12</td>
<td>9</td>
<td>-0.42</td>
<td>0.9199</td>
</tr>
</tbody>
</table>

age comparison with general study population

<table>
<thead>
<tr>
<th></th>
<th>multiple reinf. pop.</th>
<th>remaining population</th>
<th>Z value of 2 tailed WRS-test</th>
<th>2-tailed significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>number of men</td>
<td>15</td>
<td>11819</td>
<td>-3.34</td>
<td>0.0008</td>
</tr>
<tr>
<td>median age</td>
<td>22.7</td>
<td>27.9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>number of women</td>
<td>4</td>
<td>10591</td>
<td>-2.1</td>
<td>0.0375</td>
</tr>
<tr>
<td>median age</td>
<td>18.5</td>
<td>24.5</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Increased test sensitivity

The data are incompatible with the null hypothesis that type of test used makes no difference in proportion of positives detected for cervical swab specimens. Cervical swab specimens of women are more likely to test positive for *Chlamydia* if an LCx test (proportion positive: 0.124) is used instead of a culture test (proportion positives: 0.061), with the ratio between proportions being 1.91 (95% CI: 1.85-1.96, tab. 12).

Table 12: Comparing LCx tests with culture tests. Results of testing the null hypothesis that proportion of positive diagnoses in cervical swab testing is the same regardless of test used (LCx, culture). Also given are the odds ratio and its confidence interval and the result for testing the null hypothesis of equal age distribution between both test groups.

<table>
<thead>
<tr>
<th></th>
<th>negative results</th>
<th>positive results</th>
<th>total</th>
<th>proportion of positives</th>
</tr>
</thead>
<tbody>
<tr>
<td>Culture</td>
<td>2175</td>
<td>134</td>
<td>2309</td>
<td>6.10%</td>
</tr>
<tr>
<td>LCx</td>
<td>1462</td>
<td>182</td>
<td>1644</td>
<td>12.40%</td>
</tr>
<tr>
<td>Total</td>
<td>3637</td>
<td>316</td>
<td>3953</td>
<td></td>
</tr>
</tbody>
</table>

Chi-square: 35.5
Significance (df1): <0.00001
Odds ratio of proportions LCx:culture (95% CI) 1.91 (1.54-2.36)

<table>
<thead>
<tr>
<th>age comparison</th>
<th>patients with culture test</th>
<th>patients with LCx test</th>
<th>Z value of 2 tailed WRS-test</th>
<th>2-tailed significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of women</td>
<td>2309</td>
<td>1644</td>
<td>-1.1</td>
<td>0.2635</td>
</tr>
<tr>
<td>median age</td>
<td>24.7</td>
<td>24.7</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
VII. Discussion

Aim of the Dissertation

*Chlamydia trachomatis* infections are a serious burden of disease in the UK (Paavonen *et al* 1996, CMO, 1997) and recently considerable resources have been directed towards a national screening pilot program (Department of Health, 2000). With a sensitive and specific diagnostic test and an efficient treatment at hand, the disease is controllable from an individual-centred medical perspective, patients “just” have to get treated. However, from a population based public health perspective, the disease has escaped control, as *Chlamydia* incidence is rising (ISD, 2000).

Some of the more serious sequelae such as ectopic pregnancy, chronic inflammation and infertility have been associated with reinfection (Hillis, 1997). Reinfection with *Chlamydia* is a sign of inadequate control measures at both the individual level, because of high-risk behaviour and inadequate health education, and at a population level, because of inefficient screening and contact tracing strategies.

Despite many studies on reinfection risk and intervals (table 1.2), some results are contradictory and others only apply to women. In addition, only one (Pimenta *et al*, 2000) out of 11 reinfection studies took place in the UK (table 1.2), yet even the most recent report on the national screening pilot study (Tobin *et al.*, 2000) leaves the question of reinfection intervals unanswered.

The availability of large hospital and laboratory based datasets covering several years allow researchers to relate clinical outcomes to risk factors and time. The primary objective of the dissertation was to give an estimation of *Chlamydia* reinfection intervals for GUM clinic patients, based on routine laboratory test results. Reinfection intervals help clinicians to decide when a patient should come back for testing and assist health economics in allocating appropriate funding for health care. Being the only GUM clinic in Lothian, the Royal Infirmary serves a population of 773,800 people (GROSa, 2000) and routine laboratory data from January 1992 to May 2000 were available for this task.

Secondary objectives were the analysis of multiple reinfections and the impact of a change in test methods on routine data. Finally, another aim of this study was to identify suitable methods applicable to routine data in order to maximise the benefit of legacy data of other health boards. If the same kind of information from other health boards were to be evaluated in a similar way, comparisons between health boards across Scotland would be possible. Any systematic differences found could suggest factors that may be implicated in infection and reinfection.
Design of the Study

Reinfection is characterised by multiple events through time, so a cohort study design had to be chosen. Cross-sectional studies only give point estimates of prevalences and associations of exposure with disease, but lack any insights into temporal associations between exposure and disease (Hennekens et al 1987). Cohort studies are observational, exposure to risk factors are recorded through time and are compared with disease-status at the end of the study. Prospective cohorts are expensive to set up and take the whole study period to complete. However, they provide strong causal evidence for links between disease and exposure, as most factors involved can be controlled for right from the beginning, and the study is “tailor-made” to answer the topic under investigation. Retrospective cohorts on the other hand are much cheaper and faster to complete, since they utilise data that had been collected already. However, many such studies rely on information that has been gathered on a routine basis, and not to answer a particular research question. Factors important to the current research project might not have been collected and the relevant data are irrecoverable since any event took place in the past and not concurrently. The people who collected the data and those who analyse them are usually different, so great care has to be taken that all important information regarding collection is communicated to those involved in analysis.

Here, the cohort consisted of the patients visiting the RIEGUM clinic between January 1992 and May 2000, exposure variables recorded were sex, age, risk group, date of test and testing history, disease (event of interest) was defined as “reinfection with Chlamydia”.

Sources of Bias

A major source of bias for retrospective cohort studies is selection bias, since entry into the study population can neither be randomized nor controlled for retrospectively and baseline characteristics of the study population might no longer be available. Considering the nature of the disease, primarily symptomatic patients will come forward for testing, and they might be physiologically different from the rest of the population. Ideally, everyone would go to the clinic, but given the stigma of a sexually acquired disease, this is not the case and a GUM patient population might differ from the normal population, e.g. in socioeconomic status, as well as in subtle characteristics of sexual attitude and behaviour. For this study, apart from sex and age, no personal characteristics were available.

38% of the Chlamydia tests done by the MML (estimation based on interval 1.5.1999 - 1.4.2000) were for GPs and not the RIEGUM clinic, so the GUM clinic study population excludes the group of patients who rather go to their GP for STD-testing. However, in Lothian, most patients testing positive for Chlamydia
are referred to the RIEGUM clinic by their GPs for treatment and contact tracing (Louise Shaw, personal communication). Contacts who are traced will be seen at the GUM clinic, however, GPs may give patients they treat a second dose of antibiotics to give to their partner, i.e. a proportion of cases might be cured without ever being tested (Dr. Sheena Sutherland, affil.). It is still important to point out that any results reported here are only applicable to GUM clinic patients.

Also, by design, only reinfection episodes within a maximum interval of 8 years could have been detected.

Another selection bias could have been introduced by excluding 227 patients because of their conflicting DOB. If these errors occurred in high-risk outreach clinic patients only, they would be underrepresented. An alternative decision-rule for excluding records with same UPI, but non-typographically different DOBs would have been to exclude those who presented during an error prone period (see below). Still another option would be the complete exclusion of these “offending” records.

Instrument bias could have been introduced by ignoring type of test for the reinfection event. By definition, all index tests had to be done before 1.1.1998 and were thus made without LCx, but 28 out of 124 reinfected patients (22.5%) had their second positive test (reinfection event) done with LCx. Depending on whom which test was done, systematic differences could have been introduced. Also, reinfection intervals would seem to be shorter because more positive cases are getting picked up (ascertainment bias).

Observer bias is unlikely to have occurred for the specimen testing, as for the actual testing, only a patient's UPI is known, neither sex nor age.

Recording bias is a problem for retrospective studies and might be introduced by a change in staff during the 8 years of study time, for example when a less experienced or new staff member is making more errors during data entry and management. There was a higher rate of conflicting DOBs within the dataset after the third quarter of 1997 (appendix, fig. 5).

Censoring bias (informative censoring) and reporting bias would have been present if a subgroup of the study population with different reinfection risk or differing personal characteristics tended to have any subsequent tests done at their GPs after their last visit at the RIEGUM clinic. Any reinfection event diagnosed by a GP would have been lost, as well as the additional time until censoring in case of a negative test.
Analysis and Interpretation of Findings

Descriptives

Descriptive analyses of the study population do not make any assumptions and do not test hypotheses, but give a general overview and serve as a valuable starting point for further “explorations” into the data. There was an almost equal number of tests done on men and women, however the age pattern varied considerably as more young women between 15 and 24 and more older men between 25-44 were tested (fig. 6.1). This might be explained with an earlier exposure of women to sexual contacts without adequate health education and persistently increased risk behaviour in a subgroup of older men. But more tests on young women and older men do not necessarily mean that these groups are more likely to test positive. However, the proportion of positives (fig. 9) is constantly higher for men older than 22. Whereas the proportion of women testing positive is falling rather steeply and reaches levels below 5% for women at age 28, men lag about 7 years behind before they reach an equally low proportion of positives (fig 9). This could indicate age-dependent differences in risk behaviour between the sexes with older men and younger women having an elevated risk of infection. However, the possibility remains that young symptomatic men and older symptomatic women refrain visiting a GUM clinic and thus escape detection. The behavioural and biological explanations for the observed differences require specific research to proof them.

Rates of reinfection within a year are between 2.0-4.8% for men and 3.2-8.8% for women. These results are in the same range as those of Pimenta et al (2000), who looked at urban STD-clinic data in England. The UK rates are considerably lower than those reported in US studies (Blythe et al, 1992, Fortenberry et al, 1999, Hillis et al, 1994), perhaps because of differences in study populations. US studies tend to include a higher proportion of young women deemed to be at high risk. It is therefore important to ensure comparability of the cultural setting of a study before any generalisations are made. Nevertheless, women aged 15-19 have the highest risk of Chlamydia reinfection within a year, which is concordant with other findings in the literature (tab. 1.1-2). This is of particular concern, since young people tend to have a reduced perception of risk and are also just beginning to explore their sexuality, yet they have more to lose in terms of sequelae.

Analysis

Any inference drawn from data can only be made at the expense of certain assumptions. These assumptions therefore require critical reflection, so that inferences are either strengthened or treated with caution.
Survival Analysis

The sub-population of the survival study is about 3 years (median age) significantly younger than the general GUM clinic population. This was to be expected, since a patient with a reinfection attended at least twice, but his or her age at first presentation was chosen, which is, by definition, the younger one. With at least 1,607 cases in each group of a Wilcoxon Rank Sum test, even small differences in median age are likely to test significant. Therefore, the test might have been inadequate to begin with.

Kaplan Meier Curves

KM estimates for median reinfection time were between 60 and 80 months for men and 54 and 77 months for women. The KM-curves did not fall steeply through y=0.5, so that the point estimates given are likely to be unreliable. In addition, the statistical measure of median survival time might be of little clinical relevance. It is hardly in the interest of public health to wait with retesting until everyone has a 50% chance of reinfection. However, a lower threshold can be chosen analogous to the graphical estimation of the median survival time. Further issues will be discussed in the outlook section of this section.

Logrank Test

Apart from the first 5 months, survival curves for men and women had a constant slope for age categories ≥15 (all), 15-19 and 20-24. For age category 25-34, the slopes were constant until t=68, for category ≥35 they were not constant at all, probably due to the low number of events for men (4) and women (1). Logrank tests were therefore reliable for age categories: all, 15-19, 20-24 and 25-34. The survival distribution between men and women was significantly different only in the 20-24 ageband. This has important consequences as it could point to different risk behaviours or exposures between the sexes who in turn would need individual intervention strategies.

Cox's Regression

Sex

In Cox1, women were at 31% lower risk of reinfection. However, not only was sex borderline non-significant (p=0.0628), the LML plots for sex intersected each other at t=16 and the assumption of proportional hazards is thus clearly violated. Great care has to be taken in interpreting model Cox1. Therefore, separate Cox's regressions for men (Cox2) and women (Cox3) became necessary. However, it could be informative to know when the hazards were not proportional any more (here: between 15 and 18 months), as this would indicate a time dependend difference in reinfection risk between men and women, which would need to be investigated further.
Prostitutes

In Cox2 and Cox3, the multiplicative hazard of $0.00002^{\text{prost}}$ is rather meaningless, because being a prostitute ($\text{prost}=1$) results in a coefficient near zero, which would nullify any influence the other covariates have and leave only the non-parametric baseline hazard function. Not being a prostitute ($\text{prost}=0$) would set the coefficient to one, i.e. no influence on hazard.

Covariates should capture variation in data in order to be of any predictive value. Although prostitutes comprise a defined risk group and the total dataset contained 721 prostitutes, only 11 fulfilled the requirements of the survival study. The imbalance in numbers between prostitutes and non-prostitutes (11:1596) reduced the statistical power considerably and made inclusion of the prost variable in regression modelling rather superfluous. This is reflected in the high p values and standard errors of the covariate (tab. 9.1-3). LML plots for prost had only one graph because all prostitutes were censored before the first reinfection event and hazard proportionality could not be tested. Including prost as an explanatory variable was certainly not a good choice.

Prior Negative Test

Having prior Chlamydia tests was included as a proxy measure for overall STD testing history, which was shown to be associated with infection risk in some studies (Hillis et al, 1994, Fortenberry et al, 1999). 197 of 1607 (12%) patients had a negative test prior to their index test and systematic differences in risk of reinfection caused by this could have been picked up. However, the p-value of the firstneg-covariate was over 0.93 for all regressions (tab. 9.1-3), which is equivalent with the statement that having negative tests prior to the index test makes no contribution beyond chance to explain variations in reinfection risks. This is also reflected in the modest increase of instantaneous reinfection risk of 1-3% in Cox1-3 (table 9.1-3). Either the true effect on reinfection risk was too small to be detected or it did not exist in the first place. Also, negative tests before 1992 were not accounted for. The LML plots show nearly parallel graphs with the lines intersecting at t=58, which could point at a violation of the proportional hazards assumption. However, at the intersection point, only 11 out of 197 patients were left which makes the plot rather unreliable at this point.

Speaking from the point of regression analysis only, previous negative tests had no predictive value for reinfection risk. From a modelling point of view, however, this is an important prerequisite for using Markov chain simulation algorithms, because testing history (past events) must not influence infection probability. More precisely, previous negative tests had no influence on repeated infection risk and for additional proof of Markov property it remains to be shown that previous positive tests had no influence either.
Age Categories

Including a categorical covariate requires a reference group for dummy-coding (tab. 8). That group should be sufficiently big in order to have enough power to detect differences. Since the regression coefficients will be estimated in relation to the reference category, its suspected effect on risk should be on either end of the scale to make interpretations easier. Here, the group with the lowest suspected risk, agecat $\geq$ 35 was chosen as reference category.

Only in Cox1 and Cox3, agecat had a significant influence on reinfection risk and then only the first category, agecat 15-29 was significant. Instantaneous risk of reinfection for both sexes (Cox1) increases by 86% (15-19), 5% (20-24) and decreases by 16% for those aged 25-34. In women (Cox3), risk increases by 90% for 15-19 old and decreases by 16% and 20% for 20-24 and 25-34 old. In Cox2 (men), agecat 20-24 had a 32% higher risk of reinfection compared with a 23% increase for 15-19 year old.

Men aged 25-34 had a 5% lower chance of reinfection. However, the risk profile for men has to be interpreted with caution, as agecat was not significant (p=0.6476).

The regression results reflect the observations made in descriptive analysis, namely that young women between 15-19 and men between 20-24 have an elevated risk of contracting Chlamydia, even if controlled for other covariates. LML plots for age showed parallel lines for agecat 15-19, 20-24 and 25-34. The line for agecat $\geq$ 35 intersected others several times, but with only 5 reinfections out of a total of 86 in that age category, any event would have caused a rather steep step in the graph, thereby intersecting other graphs. The LML plot for agecat $\geq$ 35 is therefore unreliable.

With no more than one significant covariate in the model, a combined LML-plot for all significant covariates as described in the methods section became unnecessary.

Multiple Visits vs. one Visit

If people think they may have contracted an STD, a visit to the GUM clinic is likely. However, different people have different visit patterns and it is important to look at systematic differences between those who come more often than others in order to be able to focus health education. Looking at the variables sex and age at first visit, men and women visiting the clinic once only are significantly older (men:28.1, Women:25.0) than those with two or more visits (men:27.4, women:23.4). It is difficult to decide whether this has clinical relevance, since the groups compared in the WRS-test had at least 2900 cases each, and small differences in age are therefore likely to be statistically significant. For men, an age difference of 8 months seems hardly relevant. The difference for women is 19 months and could point at a systematic difference between women who come more than once and women who do not.
Multiple Reinfections

It does not seem to take longer for a second reinfection than it took for the first which supports the view that risk behaviour of individuals stays constant, otherwise intervals would shorten or become prolonged. Patients with multiple reinfections were significantly younger than those with only one (tab. 11). This was expected because any patient with recurrent episodes needed to be in the study for a longer time and was hence younger at his or her first visit. Looking at age to check for differences between groups does not seem to be a good diagnostic test. Also, comparing multiple reinfection episodes within 8 years automatically selects those patients with short intervals. If they were systematically different, it would have introduced bias.

Diagnostic Test Performance

Cervical swabs of women visiting the clinic between September 1997 and August 1999 were almost twice as likely to test positive if diagnosed with LCx instead of culture media (tab. ###). This means that prior to LCx testing, a proportion of people with Chlamydia were told they were not infected, did not get treated and had most likely spread the infection. Future studies on Chlamydia risk factors have to consider type of diagnostic test as a confounder and thus should include it as a covariate. Also, higher incidence rates might merely reflect the improved sensitivity of laboratory tests. Therefore, nationwide statistics on Chlamydia incidence such as those published by ISD (ISD, 2000) should include the proportion of laboratories using DNA-amplification tests. A conversion factor between old (pre-LCx) and new (LCx) rates would be:

\[
\text{incidence}_{\text{adjusted}} = \text{incidence}(\text{new}) \times (1 - \text{proportion of LCx tests}/1.91)
\]

odds ratio for proportion of positives LCx:culture = 1.91

As of 1999, 6% of GUM clinics in the UK were using DNA amplification assays (David, 1999). Ideally, both tests, LCx and culture would have been carried out here on the same individuals and compared against each other with McNemar's chi-square test, addressing the number of discordant pairs.

Here, an ordinary chi-square test was performed and the age of both sub-populations (LCx and culture) was compared with a WRS test. Median age was 24.7 for both groups and the data supported the null hypothesis of no difference in age (p=0.2635, WRS). Yet, an unknown confounder could have lead to a low risk group of women visiting the clinic between September 1997 and August 1998. Alternatively, a high-risk group of women could have begun visiting the clinic from the first of September 1998. The latter two scenarios seem rather unlikely, however, and homogeneity of both groups, “LCx-women” and “culture-women” is assumed.
The above example on test sensitivity illustrates the differences between prospective and retrospective cohorts. A proper prospective cohort study would have conducted both tests on the same individuals or, for a certain time, would have selected individuals at random and test them with either the old or the new test. In a laboratory with a high volume of tests such as the MML, this extra burden would have required additional personnel and material, which in turn would have increased the total budget of the study. Retrospective studies, on the other hand, have to find ways to minimize the impact of confounders. Here, tests were stratified for sex and specimen and only test results of cervical swabs within a certain time were compared. This reduced temporal effects and removed selection bias based on sex, since asymptomatic men are more likely to get tested with LCx than with culture. Further, age comparison between women of both groups demonstrated a certain degree of homogeneity in the retrospective cohort.

Limitations of this Study

The value of any conclusions in an observational study depends on the comprehensiveness of potentially relevant factors considered (Bull et al., 1997). The individual reasons to visit a GUM clinic are manifold, among them recent exposure to infection, presence of symptoms or general anxiousness and health concerns. This study for reinfection intervals makes two major assumptions about the comprehensiveness of the data: first, anyone with a Chlamydia infection in Lothian gets tested and second, if they do, they come to the RIEGUM clinic as a monopolist provider for all their tests. The first assumption is very optimistic, because some people are too afraid to see a doctor for STDs and never go to a clinic. Also, Chlamydia infection is asymptomatic in 50% of men and 70% of women (CMO, 1997) and these people will not feel ill so will not attend for tests unless identified as a contact and then they may be very reluctant to come for testing. A certain amount of these asymptomatics can be picked up through contact tracing or, if they are female through routine health visits such as cervical smear testing. However, cervical screening is instigated only if over 25 or over 20 if sexually active, so the major pool of infectious women under 20 would not get screened. There will still be a large amount of asymptomatic carriers in the general population, which remain undetected by looking at routine data only.

The second assumption of a monopolist provider is implicitly made during estimation of reinfection intervals, which is based on GUM clinic visit history. Chlamydia tests done outside the GUM clinic are not accounted for in the study and will lower the accuracy of estimating reinfection intervals. This is particularly worrying if one considers that e.g. between 1.5.1999 and 1.4.2000, 3528 women were tested at the GUM-clinic, compared to almost 4588 that were tested outside the GUM-clinic. Therefore, including and linking GP data would greatly enhance any future study.
Another limitation of this study is that despite using data from 8.4 years, year of index case was not included as a covariate. It would have required six additional nominal variables, which in turn would have reduced the power of the analysis considerably. However, no major STD prevention campaign took place in Lothian during the study period (Gordon Scott, personal communication), which could have influenced risk behaviour. The only known time-dependent confounder was the change in testing methods in September 1998. By definition, no index test was after 31.12.1997. However, 22.5% of reinfected patients had their second positive test (reinfection event) done with LCx and the influence of year of event (reinfection) happening could have been tested by including “test type of event” as a time dependent covariate. Also, if variation within a group is not accounted for by a covariate and if that variation is relatively large compared to between group variation, the estimation of risk will become more imprecise. The large time interval could have also lead to substantial differences in the composition of the study populations between 1992 and 1999/2000. However, Scotland has low immigration and emigration rates between 1-2% (Grosb, 2000), so this seems to be minor issue.

Age categories were chosen according to those used by ISD (2000). They might have been to coarse, so that important age effects were “diluted” during the analysis.

KM curves only account for one factor, whereas Cox's regression analysis allows adjustment for multiple explanatory factors. However, in this study the covariates available for analysis were rather unsatisfactory and the models had severe limitations. One covariate, sex, violated the assumption of proportional hazards and the regression had to be made for each sex separately, reducing the overall power. Of the remaining three covariates firstneg, prost and agecat, only in women, one (agecat) was significant and even then, only one category (15-19) had a p-value below 5%. With so many non-significant covariates the value of using Cox's regression analysis is questionable. Therefore, a purely descriptive Kaplan Meier analysis for men and women with age as factor would have been sufficient.

Even then, the gradient of the KM survival curves was so small, that any point estimate given for median time to reinfection is rather vague and has to be interpreted with great care. This is unsatisfactory, since a major goal of the study was to estimate this very interval. It can be seen qualitatively, however, that young women aged 15-19 have the shortest median reinfection interval and men aged 25-34 have the longest. A previous study by Hillis et al (1994) showed that within 5 years, 54% of women under 14 (at index case) got reinfected with Chlamydia. The median reinfection interval, i.e. the time by which 50% of the sample had become reinfected, for 15-19 year old women was 4.5 years, which compares to the result of Hillis et al (1994).
A general limitation of the Cox's regressions presented here was that 760 out of 1607 cases were not part of the “risk set” at the time of event anymore. In other words, they were censored before the first event happened and were therefore excluded during model building, which in turn reduced the power of the parameter estimation in regression modelling.

Outlook and further Research Strategy

Arguably, the greatest weakness of this retrospective cohort study was the lack of important observational variables such as socioeconomic status, occupational class, ethnicity, personal risk behaviour and STD infections other than *Chlamydia*. Given extra time, additional information on postcode, occupational class, ethnicity, number of regular/irregular sex partners, contraception used and non-*Chlamydial* STDs could have been obtained from the RIEGUM clinic and the MML if more time were available. Including both, ethnicity and socioeconomic status would help to disentangle their effects on each other.

Discussions are under way to link some of this data to the reinfection database so far constructed. Also, any covariate used in the regression should capture enough variation and have sufficient events to be of analytical value. Age categories in particular were very unevenly distributed (tab. 8). An alternative strategy for setting age categories could be to look at data from one year only, take cut-off points that capture differences in proportion of positives and then exclude this year in the analyses to avoid biased significance tests.

Subsequent reinfections could have been analysed analogously to first reinfections. Of all patients with a reinfection (124), those with at least 3 tests would be included in the study (117). The second positive test (reinfection event of first survival study) would mark the starting point of observation and the third positive test would be the “(multiple) reinfection event“ (21 patients). However, a better way of evaluating multiple reinfections would be time-to-event models that allow for multiple events (Clayton, 1994).

Coefficients of significant covariates that have a strong effect on risk in Cox's regressions could be used to create a prognostic index. The index could be subdivided into low, medium and high-risk categories. A KM plot stratified for these categories could then serve as a descriptive, graphical decision support, since retest intervals could be chosen flexibly depending on the tolerated reinfection probability.

New approaches for screening such as home sampling and reinforced contact tracing efforts have be considered to increase “catchment” of asymptomatic carriers and, ultimately, lower the prevalence of *Chlamydia* and other STDs.
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There was not enough time to go into detailed reinfection modelling. However, results from the descriptive analysis would have been combined with information from published cross sectional studies as follows:

- conceptions, abortions and teenage pregnancies (ONS, 1999, Scottish Executive, 1999a)
- attitudes towards sexual relations (NSR, 1998, Scottish Executive, 1999b)
- number of sexual partners (ONSHEA, 1998)
- income distribution (DSS, 1998)
- sexual behaviour of young people (Scottish Executive, 1999a)
- attendance at family clinics (Scottish Executive, 1999a)
- sex education (Scottish Executive, 1999b)
- STD incidence (ISD, 2000)
- social deprivation (Scottish Executive, 1999a)

The model would have followed the approach of Kretzschmar *et al* (1996) who studied the spread of STDs within a population, taking into account the structure of sexual contact patterns and different prevention strategies such as screening of subgroups, contact tracing and condom use as a crucial aspect of sexual risk behaviour. Modelling would have been done with Markov Chain Monte Carlo (MCMC) methods using WinBugs for simulations (Gilks, *et al*, 1996, BUC/DEPICL, 2000).

On a more philosophical note, access to cleaned, anonymised raw epidemiological data such as that extracted for this study could be transferred to an internet based “Open-Source” epidemiological community. Ownership of the datasets would remain with their originating institutions, but everyone would have free access to them and any changes or novel evaluation methods would have to be made electronically accessible for free as well, including custom software or computer macros for SPSS, SAS, S or other statistical packages. Patient identifiable information would have to be treated with greatest care according to the guidelines set forth by the Caldicott Committee (NHS Executive, 1999, 2000, tab. 13, appendix). The “Open-Source” idea will be discussed with the owners of this dataset.

In computer software engineering, the “Open-Source” idea has led to a proliferation of free, highest quality software, some of which is responsible for 60% of all internet services worldwide. Free flow of high quality epidemiological data would lead to a substantial increase of interdisciplinary cooperations across the globe between experts of different fields such as mathematical modelling or disease surveillance.

Knowledge would increase exponentially and lead to novel approaches that help to remove the “spirit of sickness” before it takes shape.
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VIII. References


Camus, M. 'Case to variable ratio in logistic regression'. mcamus@videotron.ca, (1.9.2000)

Clayton, D. G. (1994). 'Some approaches to the analysis of recurrent event data'. *Statistical Methods on Medical Research*, 3, 244-262
Estimating Reinfeciton Intervals for *Chlamydia trachomatis*

Clinical Effectiveness Group (CEG) (1999). 'National guideline for the management of Chlamydia trachomatis genital tract infection'. *Sexually Transmitted Infections*, 75 (Suppl), S4-S8


Diekmann, O., Heesterbeek, J. A. P. (2000). 'Mathematical Epidemiology of Infectious Diseases'. Chichester, John Wiley & Sons


Estimating Reinfection Intervals for *Chlamydia trachomatis*


Kjær, H. O., Dimcevski, G., Hoff, G., Olesen, F., Østergaard, L. (2000). 'Recurrence of urogenital Chlamydia trachomatis infection evaluated by mailed samples obtained at home: 24 weeks’ prospective follow up study'. *Sexually Transmitted Infections*, 76, pp169-172

Estimating Reinfecion Intervals for *Chlamydia trachomatis*


Paavonen, J. (1997). 'Is screening for Chlamydia trachomatis infection cost effective?'. *Genitourinary Medicine, 73*, pp 103 – 104


Estimating Reinfecion Intervals for *Chlamydia trachomatis*


Estimating Reinfection Intervals for *Chlamydia trachomatis*

SIGN (2000). 'Management of Genital Chlamydia trachomatis Infection'. Scottish Intercollegiate Guidelines Network,


Volterra, V. (1926). 'Fluctuations in the abundance of a species considered mathematically'. *Nature*, 118, S.558-560

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IX. Appendix