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The incidence of primary glomerulonephritis worldwide: a systematic review of the literature

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SHORT SUMMARY

A systematic literature review of the incidence of primary glomerulonephritis is presented. 40 papers published between 1980 and 2010 were critically appraised and included in the review. Overall, incidence was found to be between 0.2 and 2.5 /100 000/year in adults with lower incidence rates in children, for most types of glomerulonephritis.

ABSTRACT

Background

Little is known about the worldwide variation in incidence of primary glomerulonephritis. The objective of this review is to critically appraise studies of incidence published 1980-2010 so that an overall view of trends of these diseases can be found. This will provide important information for determining changes in rates and understanding variations between countries.

Methods

All relevant papers found through searches of Medline, Embase and ScienceDirect were critically appraised and an assessment was made of the reliability of the reported incidence data.

Results

This review includes 40 studies of incidence of primary glomerulonephritis from Europe, North and South America, Canada, Australasia and the Middle East. Rates for the individual types of disease were found to be in adults, 0.2/100 000/year for membrano-proliferative glomerulonephritis, 0.2/100 000/year for mesangio-proliferative glomerulonephritis, 0.6/100 000/year for minimal change disease, 0.8/100 000/year for focal segmental glomerulosclerosis, 1.2/100 000/year for membranous nephropathy and 2.5/100 000/year for IgA nephropathy. Rates were lower in children at around 0.1/100 000/year with the exception of minimal change disease where incidence was reported to be 2.0/100 000/year in Caucasian children with higher rates in Arabian children (9.2/100 000/year) and Asian children (6.2-15.6/100 000/year).

Conclusions

This study found that incidence rates of primary glomerulonephritis vary between 0.2/100 000/year and 2.5/100 000/year. The incidence of IgA nephropathy is at least 2.5/100 000/year in adults: this disease can exist subclinically and is therefore only detected by chance in some patients. In addition, referral policies for diagnostic biopsy vary between countries. This will affect the incidence rates found.

INTRODUCTION

Although much is known about clinical characteristics and natural history of the primary glomerulopathies, very little information on the epidemiology of these diseases is available from reviews. Insight into the baseline incidence of glomerulonephritis throughout the world can provide important information on trends of disease occurrence by sex, age and geographical location. New vaccines are being introduced and concerns have been raised about the potential associated risk of autoimmune diseases (46,47). It is therefore of interest to know what the baseline incidence rates across the world are so that concerns about possibly associated increased incidence rates of autoimmune diseases, such as glomerulonephritis, can be evaluated.

To our knowledge no other systematic review of incidence of the most common of the primary glomerulopathies has been conducted in the last three decades. In this paper we perform a systematic review, critically appraising studies of incidence of primary glomerulonephritis throughout the world.

Method

Searches of the Medline, EMBASE and Science Direct databases (1980 – June 2007) were carried out using the search terms 'glomerulonephritis,' 'IgA nephropathy', 'membranous nephropathy', 'membranoproliferative glomerulonephritis', 'mesangial proliferative glomerulonephritis', 'minimal change disease', 'focal segmental glomerulosclerosis', 'postinfectious glomerulonephritis', 'idiopathic crescentic proliferative glomerulonephritis', 'ANCA-associated necrotising crescentic glomerulonephritis', 'antiglomerular basement membrane disease', 'kidney disease', 'incidence', 'incid*' and 'epidemiology.' In Medline, the individual disease names were searched for as well as the term 'glomerulonephritis' because this MeSH term does not include all types of glomerulonephritis as daughter terms in its hierarchical structure. The inclusion criteria were that the studies reported original work, that the study reported incidence of specific forms of glomerulonephritis with reference to a denominator population, that the estimates of population size and person-time contributed were accurate and that efforts had been made to ascertain all incident cases. When assessing the likelihood of missing incident cases, papers were evaluated as follows: 1) for case finding studies, did the authors ensure that all of the subjects contributing to incidence denominator data would have been eligible to have the disease diagnosed and did the authors check all relevant medical records? 2) For all studies, were cases checked to ensure that they were incident and not prevalent? 3) For all studies, did the authors ensure that they cause of glomerulonephritis was autoimmune and not secondary to another disease? Where possible we only included incidence rates for cases of glomerulonephritis caused by autoimmunity, determination of which relied on information given in the paper.

The titles and abstracts of all of the studies produced by the searches were reviewed and those papers accepted for inclusion in the study were appraised. Studies published in English, French, German, Spanish or Dutch were included. Review papers identified were searched for secondary references reporting on original research; secondary references found from any of the other papers reviewed were also included.

A standard data abstraction form was used to record all details of the papers reviewed; a copy of this is given in the Appendix, figure 1A. Each study was scored for accuracy of the incidence rates it presented and was classified as being at low, medium or high risk for under- or overestimation of reported incidence rates by considering the reliability of numerator and denominator data. For instance, inclusion of prevalent cases or those thought not to be caused by autoimmunity will have led to overestimated rates as will underestimated denominator data. Conversely, missing cases or an overestimated denominator (e.g. a catchment area from which not all inhabitants had access to hospital services) would be considered to have resulted in underestimated incidence rates. Explanations provided by the papers' authors as to why incidence rates were as expected or whether they were considered to be an over- or underestimate of the true incidence rate were taken into account in this process. If the extent of likely error was considered to be very great, the study was excluded. To minimise subjectivity, this assessment was agreed between two of the authors and random checks were performed to ensure consistency. Rates are presented as the number of cases/100 000/year and where sufficient data were given in the paper, rates were checked for accuracy. Guidelines were followed in the reporting of this study to ensure that key information was presented (1).

Results

The results of the database searches with the number of included and excluded papers are given in figure 1; the excluded references are available on request from the authors.

Most of the papers rejected at abstract review stage did not report on primary and autoimmune glomerulonephritis and had been found from the search using 'kidney disease' as the search term. Of the remaining papers, reasons for rejecting included those reporting on an ill sub-group of the population (e.g. those with systemic lupus erythematosus), those that reported on prevalence and not incidence, review papers and those that gave incidence rates as a percentage of people who had a renal biopsy.

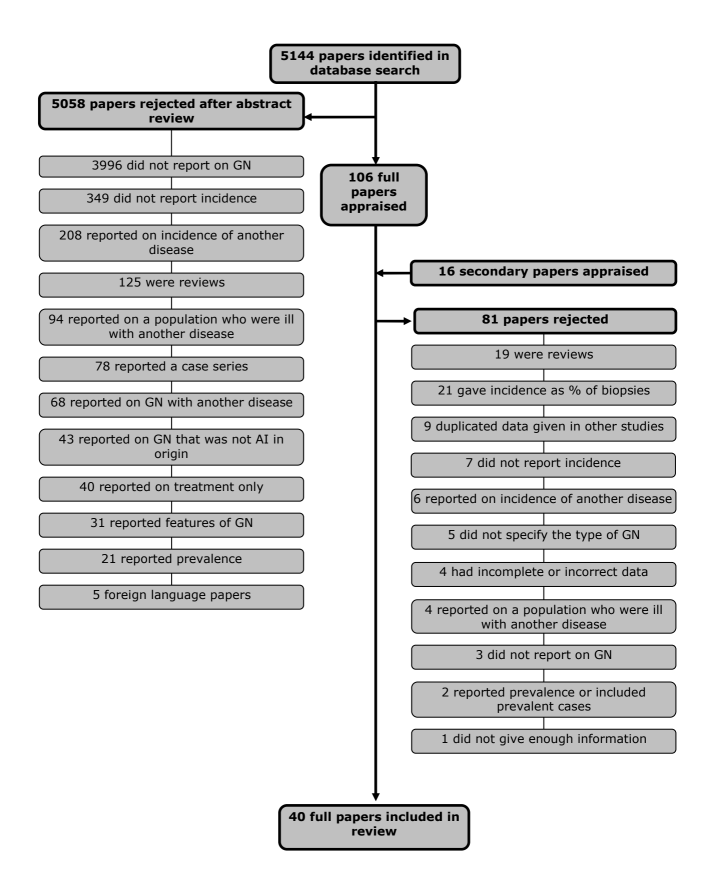


Figure 1: Results of the database searches showing the number of references found, those excluded and the final number of papers included in this review. (GN = glomerulonephritis; AI = autoimmune)

For some types of glomerulonephritis, the terms used by authors varied, some using the term to describe clinical presentation or syndrome for example crescentic proliferative (2) or crescentic glomerulonephritis (3-5), nephrosis (2,6), rapidly progressive glomerulonephritis (7,8), acute glomerulonephritis (9) and acute nephritis (10) whereas others used non-specific terms such as vasculitis (11-15) and total glomerulonephritis (9). These incidence rates have been excluded from the review because it was impossible to determine the extent to which these papers reported on primary and autoimmune glomerulonephritis.

Descriptions of the studies are given in table 1 and incidence rates are displayed in forest plots showing rates for children in figure 2 and adults or all ages in figure 3. Appendix 1, table 1A includes a table of all rates presented in the forest plots. Most of the studies included in this review investigated populations in Australasia (3,11,16), Europe (2,4-6,9,14,17-30), and North America (31-36) with five studies from the Middle East (13,37-40), two studies from South America (15,41), one study from Japan (42) and one study from Tunisia (43). Findings for the different subtypes of glomerulonephritis are summarised below.

IgA nephropathy: four studies reported rates in children and teenagers: 0.03 /100 000/year (CI₉₅ 0-0.1) in Venezuela (41), 0.08 (CI₉₅ 0.01-0.46) /100 000/ year (1975-1984) to 0.57 (CI₉₅ 0.23-1.18) /100 000/ year (1985-1994) (0-17 years) in Tennessee, US (33), 4.5 /100 000/year (0-15 years) in Japan (42) and 0.31 /100 000/year (0-15 years) in Italy (25) (CI₉₅ indicates 95% confidence interval). The difference in these rates may be explained by the study type: the Japanese study was a screening study and therefore detected subclinical cases of IgA nephropathy as well as including three possibly prevalent cases (total 37) who did not have previous screening results recorded. In contrast, the study from Tennessee and Italy were based on renal biopsy and the study from Venezuela used chart reviews of symptoms and some biopsy results.

Most of the other studies were prospective, reporting rates from 0.2 /100 000 /year to 2.8 /100 000/year (2-6,13,18,19,21,27,43); the retrospective studies reported a similar range of rates, from 0.4 /100 000/year to 2.9 /100 000/year (9,11,14,20,29,30,35,36,38,39). Two studies reported incidence rates of 5.0/100 000/year (28) and 5.7/100 000 /year in males (11) but both had high biopsy rates which is likely to have contributed to the greater incidence rates found. Many of the studies included in this review were conducted between 1970 and 1990; no trend in incidence rates with time was discernable. Sixteen studies (3-6,9,11,13,14,20,21,25,27,29,32,35,38) reporting rates used immunofluorescence in diagnosing IgA nephropathy.

Membranous nephropathy: two studies (25,31) gave incidence rates in children and adolescents: 0.05 /100 000/year (1985-1993) to 0.09 /100 000/year (1993-2002) (6 months – 19 years) in Ottawa-Hull Canada (31) and 0.02 /100 000/year (0-15 years) in Italy (25). For the other studies, most were prospective and reported incidence rates between 0.3/100 000 /year and 1.4 /100 000/ year (2-6,13,18,19,21,27,28); the retrospective studies reported incidence to be between 0.2/100 000/year and 1.3/100 000/year (9,11,14,15,20,26,29,30,35,38,39,44). Not many studies reported on differences in rates between males and females. In those that did, the numbers were low and no confidence intervals or indications of statistical significance for differences in incidence rates were available. Three studies (6,9,11) suggested the rates were higher in males than females whereas El Reshaid et al. (13) reported the opposite. Taking into account the methods used in the studies presented, it is not thought that there has been a change in incidence between 1970 and 2000. The studies that used retrospectively collected data are likely to have missed cases and therefore underestimate the incidence rates. There was insufficient information to conclude reliably whether there is a difference in risk between males and females. Our best estimate of the incidence is 1.2 /100 000/year.

Membrano-proliferative glomerulonephritis: thirteen studies were retrospective and six were prospective; the range of incidence rates in all

studies was between 0.14 /100 000/year and 0.93 /100 000/year. Over time, incidence appears to have decreased from around 0.7 /100 000/year in the 1970s (3,6,18) to 0.2 in the 1990s (2,5,9,14) with the exception of Covic et al (26) who report a rate of 0.93/100 000/year in 2004. Covic *et al* (26) link this to the higher rates of streptococcal infection and hepatitis B and C in their population; they also note a decrease in the prevalence of membranoproliferative glomerulonephritis between 1995 and 2004 which they believe is associated with improvements in income, sanitation, social and medical infrastructure. Simon et al. (2,6) noted that there was an association between streptococcal infection and the onset of membrano-proliferative glomerulonephritis and that the decreases in incidence rates of membrano-proliferative and post-streptococcal glomerulonephritis were closely linked. Other differences in rates are noticeable: Hachicha et al. (43) found the highest rate of membrano-proliferative glomerulonephritis although this is probably an over-estimate of the true rate; in New Zealand (3) an incidence rate five times higher in Polynesians than in non-Polynesians was reported. This may indicate an infection related glomerular-specific injury. Two studies (2,3) reported on the incidence of type I membrano-proliferative glomerulonephritis, two studies (9,14) reported that 68% of their cases were type I and in a third study (30) 90% were type I; the remaining studies (3,5,6,11,18,19,25-27,31,38,39,45) did not specify the type of membrano-proliferative glomerulonephritis diagnosed.

Mesangial proliferative glomerulonephritis : four studies included were prospective and three were retrospective; all gave incidence rates between 0.2 /100 000/year and 1.1/100 000/year (3,5,6,26,27,44). In order to differentiate mesangial proliferative glomerulonephritis from IgA nephropathy, immunofluorescence should be used. All seven studies used immunofluorescence in their studies but four reported that this was not performed on all biopsies (26,27,30,44) giving a potential overestimation of the incidence rate. The only study to make the distinction between IgA nephropathy and mesangial proliferative glomerulonephritis without IgA deposits was that by Schena *et al* (5), who found an incidence rate of

0.16/100,000/year therefore it is likely the 'true' incidence rate of mesangial proliferative glomerulonephritis is at the lower end of the range given.

Minimal-change disease: in children, minimal change disease has been found to cause over 75% of cases of nephrotic syndrome (46). Seven studies reported on incidence of nephrotic syndrome, which would produce a slight overestimation of incidence of minimal change disease (22,24,34,37,40,41); three studies reported on incidence of minimal change nephrotic syndrome (23,25,31). Incidence rates in children were between 0.23 /100 000/year and 15.6 /100 000/year (22-25,31,34,37,40,41). The rates were reported with respect to ethnic origin and differences were noted: 0.23 -2.8/100 000/year in Caucasian children (22,24,25,31); 2.4/100 000/year in Hispanic children (41); 3.4 /100 000/year in Afro-Caribbean children (22); 7.2-11.6/100 000/year in Arabian children (37,40) and 6.2-15.6/100 000/year in Asian children who resided in the UK (22,23).

In the remaining studies, retrospective and prospective studies reported similar rates between 0.2/100 000/year and 0.8/100 000/year in adults (3,5,9,11,15,18,21,26,27,29,30,35,38,39,44); no trend of changes over time was found. Taking into account the accuracy of these rates, our best estimate of incidence of minimal change disease in adults is 0.6/100 000/year.

Focal segmental glomerulosclerosis: most rates, whether from prospective or retrospective studies, were between 0.2 /100 000/year and 1.1 /100 000/year; the Australian study reported the highest rates of 2.5/100 000/year in males and 1.8/100 000/year in females. The latter may be due to the fact that in Australia, people are referred for biopsy more often than in other countries included in this review (11). In the three studies that investigated differences in rates between males and females, incidence appeared to be higher in males (9,11,13). However no indication was given of the statistical significance of these differences and

given the fact that the numbers of cases were low, they may have arisen by chance.

Other types of glomerulonephritis: Incidence rates were presented for other types of glomerulonephritis, however these were limited to just one or two studies per disease type. Details are given in the forest plots: generally these rates were low at less than 1.0 /100 000/year. However Becquet *et al.* (16) in their study of post-infectious glomerulonephritis in French Polynesia, reported an incidence rate of 18/100 000/year; this is likely to be an underestimate of the true rate. This country had a higher rate of bacterial infections due to the climatic conditions, greater numbers of people sharing residences, low socioeconomic level and a lower use of medical care due to cultural beliefs; these factors are all thought to contribute to this higher incidence rate.

Accuracy of incidence rates

There are a number of components to assess when considering the incidence rates presented. Indication of likely accuracy of rates has been given in table 1. Of key importance is whether the cases included in the numerator were new cases: six studies reported in their methods section that only new cases were included (2,4,6,9,22,24,36,37). As glomerulonephritis is diagnosed by biopsy, the more liberal the biopsy policy, the greater the possibility of detecting all cases of the disease. Some studies gave their biopsy rate per head of population: these are given in table 1 and vary between 1.08 and 24.7 /100 000/year (6,11,14,18,19,21,26-30,35,38,39,44).

Discussion

This literature review found incidence rates for different types of primary autoimmune glomerulonephritis to be between 0.2/100 000/year and 2.5/100 000/year in adults. Most studies were from the US and France therefore it is difficult to draw conclusions regarding variability of rates with geographical location or ethnicity. Given that glomerulonephritis can exist subclinically, and given differences in access to renal biopsy between different healthcare systems, it is likely that geographical variations in incidence rates found can be explained by differences in diagnosis rather than by genuine difference in disease frequency.

It is useful to note the type of studies undertaken to determine incidence of glomerulonephritis: in reviews of incidence of other autoimmune diseases (47,48), prospective studies are thought to have given more accurate rates than retrospective studies. However, in the studies presented here the rates reported were consistent irrespective of the study type.

Most studies used biopsy to diagnose the disease for the majority of cases however only five studies (3,21,27,36,40) reported the guidelines used for diagnosis. Classification of the disease and detection threshold of signs and symptoms may cause variation in incidence rates between studies (11,14,21) and the lack of a central histopathology review may have caused within-study variations (14). Wyatt *et al.* (32) reported that over time there was an increased recognition of IgA nephropathy therefore the incidence rates from the end of the study may be more accurate. Similarly Mazzuchi *et al* (15) reported rates using a national registry and found an increase in incidence with time they reported to be due to a greater awareness of the disease and earlier diagnosis.

Cases of IgA nephropathy can exist subclinically and therefore will only be diagnosed through routine urinary tests or if a patient presents with severe symptoms (27): Simon *et al.* (6) reported that 60% of cases of idiopathic IgA nephropathy were discovered by chance through routine testing as part of a medical examination in employment. Screening populations for conditions that can exist subclinically will produce higher and more accurate incidence rates especially when done routinely so that prevalent cases are not included in the incidence rate. In this review one study (42) used a regular screening program to find cases and the rate produced was greater by nearly an order of magnitude than other comparable incidence rates. Sehic *et al.* (33) reported that many cases of IgA are never diagnosed and that limited access to medical care for those

of lower socio-economic status may explain some failure to diagnose in this study.

There is the possibility of cases of secondary glomerulonephritis being included as primary cases particularly in retrospective studies; this would lead to an overestimation of incidence rates. There are links, for example between membrano-proliferative glomerulonephritis and hepatitis B or hepatitis C. Variations between countries in terms of rates of infections will contribute to differences in incidence rates.

One of the key factors in explaining differences in incidence rates of glomerulonephritis is the difference in referral and biopsy policies between different countries and even between regions of countries (14,21,28,29). Briganti et al. (11) found higher incidence rates of minimal change disease and focal segmental glomerulosclerosis than other studies. They claim this may be as a result of a more liberal biopsy policy, leading to the detection of less well-defined or asymptomatic cases. In contrast, The New Zealand Glomerulonephritis Study (3) noted that children and teenagers with postinfectious GN or minimal change nephropathy are rarely biopsied and older patients may also be less likely to have biopsies taken than younger adults. There is evidence that Polynesians have poorer access to healthcare resources in New Zealand than others and in this study, a high proportion of Polynesian patients presented for the first time with endstage renal failure, making it impossible to reach a specific diagnosis. Lack of access to healthcare may also have resulted in prevalent cases being included in the incidence rates for glomerulonephritis in this population. Naumovic et al (30) reported that their low biopsy rate (1.08/100 000/year) may be due to economic sanctions and that elderly people and those with diabetes do not routinely undergo biopsy._Incidence was found to be lower in African-American children compared to Caucasian children (33). The study authors suggested this may be because the decision to perform a biopsy for those suspected to have mild disease was referred to their parents who may have been less likely than Caucasian parents to give their consent; records were not available for the number of biopsies refused (17).

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Renal biopsy policy also affects diagnosis rates in the elderly: a number of studies reported increases in the number of elderly patients biopsied as a result of a change in policy and increasing age of the population (2,4,9,32) with Stratta *et al.* (9) reporting that rates of elderly patients diagnosed increased from 1.6% in 1970 to 20.4% in 1994. Wyatt *et al.* reported an increase in incidence at 45 years and over which was thought to be due to a more proactive attitude towards conducting diagnostic procedures in the elderly. It is likely that incidence in older people is underestimated as not all cases were referred to specialists or underwent biopsies (9).

Conclusion

Reported incidence rates of glomerulonephritis in adults varied between 0.2 and 2.5 /100 000 /year depending on the type of glomerulonephritis. Incidence in children was generally lower with most rates around 0.1 /100 000 /year; two exceptions to this were that incidence of minimal change disease in children was around 2.0 /100 000/year and a screening study that reported a rate of IgA nephropathy of 4.5 /100 000 /year in children in Japan.

The reported incidence rates are likely to underestimate true rates of IgA nephropathy as this disease can exist subclinically and may never be detected however other types of glomerulonephritis may be overestimated due to relapses and prevalent cases being counted as incident. There is variation in biopsy policy between countries, which affects the incidence rates found. Incidence in older people appears to have increased over time: this is considered to be due to greater inclusion of this age group in referrals for biopsy rather than due to a genuine increase in disease occurrence.

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Table 1: Details of diagnosis of cases and population covered for each of the studies included in this report

Study	Cases	RUE	ROE	Biopsy	Description
				rate	
Hachicha <i>et al</i> . (43)	611	**	***	Not	Case definition: >14 years. Renal biopsy in 230 cases; histological disease type
Study type: Retrospective				given	determined. Very few study details given. Case identification: No details given.
hospital record review					
Location: Sfax, Tunisia					
Utsunomiya <i>et al</i> . (42)	37	**	**	99.5%	Case definition: Initial screening of first urine of morning; second screening of
Study type: Prospective					those with proteinuria, occult blood and urinary sediment. Those with glucosuria
screening study					were referred. Follow-up for a few months. Case identification: All students aged
Location: Yonago City,					6-15 years screened annually. Indication for renal biopsy determined according to
Japan					criteria. Records of previous screenings checked to identify new cases.
Briganti <i>et al</i> . (11)	1147	*	*	21.5	Case definition: Renal biopsy. Case identification: Retrospective review of
Study type: Retrospective					pathology reports of all biopsies; evaluated by light microscopy and
Location: Victoria, Australia					immunofluorescence or immunohistochemistry; majority assessed by electron
					microscopy. No details available of clinical findings or indication for biopsy.
The New Zealand	803	***	*	`Liberal'	Case definition: > 14 years. Renal biopsy material examined using light,
Glomerulonephritis Study (3)					immunofluorescence and electron microscopy. Uniform histological classification
Study type: Prospective					agreed; all renal biopsies re-evaluated independently using Churg and Sobin
Location: New Zealand					classification. Case identification: 4 nephrology centres covering 75% of the
					Polynesian and 84% of the non-Polynesian populations used.

Study	Cases	RUE	ROE	Biopsy	Description
				rate	
Becquet <i>et al</i> . (16)	12	*	*	Not	Case definition: < 15 years. Diagnostic criteria were microscopic or macroscopic
Study type: Retrospective				given	haematuria, decreased C3 fraction of the complement evidence of recent
Location: French Polynesia					streptococcal infection established by presence of elevated anti-streptococcal
					antibody titres. Case identification: Admitted to one hospital.
Frimat <i>et al</i> . (17)	Not	*	*	Not	Case definition: Prospective longitudinal cohort study; >15 years. Diagnoses
Study type: Prospective	given			given	based on renal biopsy and checked by pathologist. Study restricted to half
Location: East France					geographical area to ensure all biopsies included. Case identification: 17 renal
					units.
Abdulmassih <i>et al</i> . (18)	266	**	*		Case definition: >15 years, diagnosed by biopsy. Case identification: All
Study type: Retrospective			′76-80	7.2	biopsies for the area examined in one lab.
Location: Picardy, France			<i>'</i> 81-85	8.6	
Berthoux <i>et al</i> . (19)	Not	*	*	13.0	Case definition: Renal biopsy. Case identification: Sent questionnaire in 1989 to
Study type: Prospective	given				all nephrology services; also provided information for denominator.
Location: Rhone-Alps,					
France					
Simon et al. (6)	480	*	*		Case definition: Renal biopsy specimens processed and stained for light
Study type: Prospective			′76-80	18.7	microscopy and immunohistory; electron microscopy not systematically performed.
Location: St. Brieuc, France			′81-85	20.1	Case identification: Biopsy performed at hospital nephrology department;
			<i>'</i> 86-90	16.2	collaborated with major medical screening institutions.
Simon et al. (4)	131			Not	Case definition: > 60 years old, otherwise, as above.
As above				given	
Simon <i>et al</i> . (2)	898			Not	Case definition and identification: As above.
As above				given	

Study	Cases	RUE	ROE	Biopsy	Description
				rate	
Simon <i>et al</i> . (45)	898			Not	Case definition and identification: As above.
As above				given	
Schena <i>et al</i> . (5)	1293	**	**	~ 4.5	Case definition: Renal biopsies mainly evaluated by light-microscopy and
Study type: Prospective					immunofluorescence; electron-microscopy used in 38% of cases. Case
Location: Italy					identification: Biopsies collected at Italian renal units.
Coppo <i>et al.</i> (25)	256	**	**	Not	Case definition: 0-15 years; renal biopsy analysed by light microscopy and
Study type: Prospective				given	immunofluorescence; electron microscopy used in 32% of cases. Case
Location: Italy					identification: Renal units where biopsies were performed.
Stratta <i>et al</i> . (9)	454	*	*	Not	Case definition: >15 years, renal biopsy taken during time period. Biopsies
Study type: Retrospective				given	underwent light microscopy and immunofluorescence; electron microscopy not
Location: City/ Province of					routinely used. Case identification: 3 nephrology centres in city (provided renal
Turin, Italy					biopsies to virtually entire area); those outside region were excluded.
Rivera <i>et al</i> . (14)	Not	***	***	4.8	Case definition: Primary glomerulonephritis classified into eight groups; criteria
Study type: Retrospective	given				not given. Case identification: Retrospective review of renal biopsies from national
Location: Spain					registry.
Grupo de Estudio de la	1471	***	*	Not	Case definition: Diagnosis established from kidney biopsies studied by light
Sociedad Espanola de	(max.)			given	microscopy and immunofluorescence. >14 years. Membranoproliferative GN was
Nefrologia (20)					classified as type I or type II Tried to identify all primary cases. Case
Study type: Retrospective					identification: 33 hospitals responded.
Location: Spain					
Wirta et al. (28)	958	**	*	UH 24.7	Case definition: Kidney biopsy; SNOMED classification Case identification:
Study type: Prospective				CH 9.1	Patients receiving a renal biopsy at the university or central hospitals.

Study	Cases	RUE	ROE	Biopsy	Description
				rate	
Location: Western Finland					
Heaf et al. (44)	1762	**	**	3.82	Case definition: Renal biopsy classified according to WHO guidelines and presence
Study type: Retrospective					of immune deposits. Case identification: Renal biopsy register.
Location: Denmark					
Tiebosch et al. (21)	129	*	*	12.6	Case definition: Renal biopsies processed and stained for light microscopy,
Study type: Prospective					transmission electron microscopy and immunohistochemistry and classified
Location: Areas surrounding					according to WHO guidelines. Case identification: Biopsies taken in direct referral
the cities of Heerlen,					hospitals for GPs.
Maastricht and Sittard, The					
Netherlands					
Hanko <i>et al</i> . (29)	907	*	*		Case definition: > 16 years; all aduct native kidney biopsies analysed. Case
Study type: Retrospective			`76-85	2.02	identification: Renal services in pathology department at city hospital.
Location: Northern Ireland,			` 86-95	3.86	
UK			′96-05	7.08	
Sharples <i>et al</i> . (22)	44	**	***	Not	Case definition: <16 years presenting with nephrotic syndrome responding to
Study type: Retrospective				given	corticosteroids. Nephrotic syndrome defined as proteinuria of at least 3+ on Albustix
Location: Birmingham, UK					testing with oedema and a plasma albumin concentration of 25 g/l or less. Steroid
					response defined as abolition of proteinuria within eight weeks of starting
					prednisolone. Case identification: Names and hospital numbers were traced for all
					admissions with the main or subsidiary diagnosis and responding to corticosteroid
					treatment.

Study	Cases	RUE	ROE	Biopsy	Description
				rate	
Feehally <i>et al</i> (23)	43	**	**	Not	Case definition: <15 years. Minimal change nephrotic syndrome defined by
Study type: Retrospective				given	complete abolition of proteinuria within four weeks in response to corticosteroids
Location: Leicestershire, UK					with no hypertension or renal impairment. Some had further information from
					biopsy material analysed by light, immunofluorescence and electron microscopy.
					Case identification. From hospital records.
McKinney et al (24)	194	***	***	Not	Case definition: 0-15 years. Nephrotic syndrome diagnosed if proteinuria was at
Study type: Retrospective				given	least 3+ on testing with albustix, with hypoalbuminaemia and oedema. Investigated
Location: Yorkshire, UK					response to corticosteroids. Case identification: Primary: listings provided by
					paediatricians. Secondary: inpatient hospital episode statistics; identified those with
					ICD9 codes 580-583 (acute/chronic GN, NS, nephritis, nephropathy) 590 (infections
					of the kidney.) Data extracted from hospital notes. Ethnic group determined by
					surname.
Rychlik <i>et al</i> (27)	1932	**	**	5.37	Case definition: Indications for biopsy differed between centres. Histological
Study type: Prospective					evaluation by light microscopy and immunofluorescence performed routinely, with
Location: Czech Republic					electron microscopy in a number of cases. Histological classification used WHO
					recommendations. Case identification: Renal biopsy records collected from renal
					units. Questionnaire used to collect relevant data.
Covic et al (26)	Not	**	**	11.3	Case definition: >18 years; diagnosed by renal biopsy. Case identification: Two
Study type: Retrospective	given				large referral centres.
Location: Moldova and					
Banat, Romania					

Study	Cases	RUE	ROE	Biopsy	Description
				rate	
Naumovic <i>et al</i> (30)	Not	***	**	1.08	Case definition: >18 years; diagnosed by renal biopsy; stained and analysed by
Study type: Retrospective	given				light microscopy. Case identification: Nephrology unit.
Location: Serbia					
Al Arrayed et al (38)	218	**	**	5.8	Case definition: Indications for biopsy: proteinuria, unexplained microscopic or
Study type: Retrospective					macroscopic haematuria, systemic disease with clinical evidence of renal
Location: Bahrain					involvement, unexplained renal impairment and renal impairment in post-transplant
					patients. Case identification: All renal biopsies, nephrectomy specimens and
					referral slides pertaining to renal disease reviewed.
Al Arrayed et al (39)	40	**	**	5.4	Case definition: As above. Case identification: As above.
As above					
El Reshaid et al (13)	315	*	***	Not	Case definition: Histological diagnosis of biopsy made on results of light
Study type: Prospective				given	microscopy, immunofluorescence and electron-microscopy in selected patients.
Location: Kuwait					Case identification: Patients screened for glomerulopathy; those meeting criteria
					referred for biopsy.
Zaki <i>et al</i> (40)	55	**	**	Not	Case definition: Patients had oedema, albuminuria and hypoalbuminaemia. Biopsy
Study type: Prospective				given	in those non-responsive to steroid medication over 4 weeks. International Society of
Location: Kuwait					Kidney Disease in Children guidelines used. Case identification: Children admitted
					to the paediatric hospital departments of in Kuwait.
Elzouki <i>et al</i> (37)	19	**	***	Not	Case definition: <15 years. Renal biopsy performed in 17 of 19 cases. Standard
Study type: Prospective				given	methods of light and electron microscopy and immunofluorescence. Case
Location: Benghazi, Libya					identification: El-Fateh Children's Hospital and clinics in the area.
Filler et al (31)	159	*	**	Not	Case definition: Diagnosis made according to the International Study of Kidney
Study type: Retrospective				given	Disease in Children criteria, verified by chart review. Case identification: All

Study	Cases	RUE	ROE	Biopsy	Description
				rate	
Location: Ottawa-Hull					inpatients and outpatients referred to nephrology services included. Hospital
region, Canada					admission database and renal biopsy records also checked.
Wyatt <i>et al</i> (32)	192	**	*	Not	Case definition: Renal biopsy using direct immunofluorescence; standard criteria.
Study type: Prospective				given	Case identification: Primary method of ascertainment by renal pathologists,
Location: Central and					hospital records also reviewed.
eastern Kentucky, US					
Swaminathan <i>et al</i> (35)	116	**	*		Case definition: Renal biopsy evaluated with light microscopy,
Study type: Retrospective			′74-`83	8.2	immunofluorescence and electron microscopy; diagnosis confirmed by renal
Location: Olmsted County,			<i>'</i> 84-`93	8.8	pathologist. Case identification: Record linkage gives details of virtually all
US			′94-`03	17.5	medical care provided.
Fischer <i>et al.</i> (36)	112	*	*	Not	Case definition: WHO classification. Case identification: Kidney biospies from
Study type: Retrospective				given	patients newly diagnosed with IgA at the university were retrieved from inhouse and
Location: New Mexico					consultation files.
Kim et al. (34)	163	**	**	Not	Case definition: Nephrotic syndrome defined as heavy proteinuria, edema and
Study type: Retrospective				given	hypoalbuminemia; some underwent renal biopsy. Case identification: Record
Location: New Orleans					review and two main referral hospitals
Sehic <i>et al</i> (33)	17	**	*	Not	Case definition: Patients < 18 years and resident in study area. Diagnosis of IgA
Study type: Retrospective				given	nephropathy made by renal biopsy.
Location: Shelby County,					
Tennessee, US					
Orta-Sibu <i>et al</i> (41)	505	**	***	Not	Case definition: <15 years; Acute GN with hematuria, edema, arterial
Study type: Retrospective				given	hypertension present. Nephrotic syndrome was diagnosed on the basis of
Location: Venezuela					proteinuria greater than 40 mg/h per m ² body surface area with or without edema,

Study	Cases	RUE	ROE	Biopsy	Description
				rate	
					hypoproteinemia and hypercholesterolemia. Case identification: Information
					obtained by contacting 17 centres with a questionnaire. Data collected by chart
					review of patients.
Mazzuchi et al. (15)	2058	***	*	Not	Case definition: Diagnosed by biopsy; defined by minimal glomerular lesions,
Study type: Prospective				given	levels of proteinuria, serum creatinine, arterial hypertension, glomerular filtration
Location: Uruguay					rate. Case identification: National registry

RUE: risk of underestimation; ROE: risk of overestimation; * low; ** medium; *** high; ESR: erythrocyte sedimentation rate; ANA: anti-nuclear antibodies; GBM: glomerular basement membrane; ELISA enzyme-linked immunosorbent assay; biopsy rate: /100 000/year

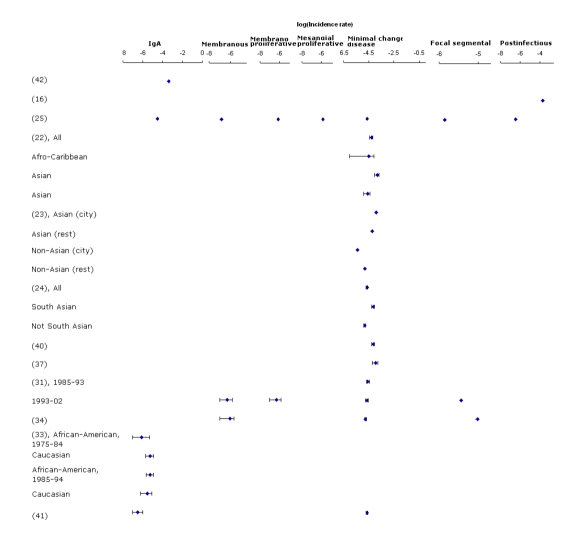


Figure 2: Incidence of glomerulonephritis in children and adolescents.

Anti-GBM Postinfectious Membrano Mesangial Minimal Focal ΙgΑ Membranous proliferative change disease segmental proliferative. 8 -6 -4 .-8 -4 -5 -5.5 -5 -6 -4 -7 -5 -6 -8 -8 -6 -7 (43), 1977-90 . . . (11), 1995, 97, Males . ٠ ٠ . (11), 1995, 97, Females ٠ ٠ ٠ ٠ (3), 1972-83, non-Polynesian ٠ ٠ ٠ ٠ . ٠ . ٠ (3), 1972-83, Polynesian ٠ . . ٠ . . (17), 1988 (19), 1987-88 . ٠ * (18), 1976-80 (18), 1981-85 M н H ⊢€I ٠ ⊢+I ⊢€I 10 ٠ (6), 1976-90, Males . (6), 1976-90, Females ٠ (4), 1976-90 (45), 1976-90 ⊢-•-+ H ٠ ٠ (6), 1976-85, 20-79 ٠ (6), 1986-95, 20-79 ٠ (6), 1996-02, ≥ 20 ٠ ٠ (6), 1976-90, 10-79 (4), 1976-90, >60 (6), 1976-90 ٠ ٠ ю ٠ **—** (6), 1976-85, 20-59 ٠ ٠ (6), 1976-85, 60-79 ٠ (6), 1986-95, 20-59 . (6), 1986-95, 60-79 (6), 1996-02, 20-59 (6), 1996-02, 60-79 ٠ (6), 1996-02, ≥ 80 ٠ (4), 1976-90 10 (5), 1993٠ ٠ H . + ٠ ٠ (25), 1992-94 (9), 1990-94 . ٠ ٠ . (14), 1994-99 ٠ н (20), 1977-86 ٠ (28), 1980-00, University hospital . (28), 1980-00, Central hospitals (44), 1985-97 ٠ -. . . (21), 1978-84 . . (29), 1976-05 ٠ н н (27), 1994-00 . H (26), 2004 (30), 1987-06 ⊢+I ٠ ٠ ٠ ٠ H (38), 1990-02 H H♦H H M ⊢+I (39), 2003-06 H€H **—** ٠ ⊢ŧ **⊢**→+ (13), 1995-01 . ٠ (32), 1975-84 (32), 1985-94 (36), 2000-05 . H (35), 1974-03 M **H** M (15), 1990-94 ٠ (15), 1995-99 ٠ .

log (incidence rate)

Figure 3: Incidence in adults or those of all ages

(15), 2000-03

Appendix

Ref Manager ID	C C	Case definition		Risk of missing cases Reasons
Year published Reviewer	0			Low Medium High
Original in English?		Number of cases	0	
Translation	~	Base population:	Base population:	Risk of overestimating cases
Excluded?		person years	number of people	
Reason for exclusion		0	U	
		Source of case identificat	ion	Secondary references
				Notes
Study dates		Country		
		Region		
Run in period		Ethnic distribution		
Ref Manager ID)	0	Type of rate	×
Race		~	Incidence	0
Other ethinc ori	gin		Units	×
			Lower Cl	0
Gender		~	Upper Cl	0
Multiple location	ns?		Figures checked?	
Time period			Figures correct?	
Other descripto	r		Notes	
Age range				

Figure 1a: Data abstraction forms

Ref	Race	Location	Time	Age	IgA	Membranous	Membranopr	Mesangial	Minimal	Focal	Anti-	Postinfecti
			period	range	nephropa	nephropathy	oliferative	proliferativ	change	segmental	GBM	ous GN
					thy		GN	e GN	disease	glomerulos	disease	
										clerosis		
(43)	All	Sfax,	1977-90	>14	0.3 (0.2,	1.1 (0.8, 1.4)§	3.1 (2.7, 3.6)		1.2 (0.9,	1.0 (0.8,		
		Tunisia		years	0.6) §		§		1.5)§	1.3)§		
(42)	(42) Asian	Yonago	1983-	<15	4.5							
		City,	1999	years								
	Јара	Japan										
(11)	Caucasian	Victoria,	1995,	All	M: 5.7, F:	M: 1.8, F: 0.8	M: 0.3, F: 0.2		M: 0.7, F:	M: 2.5, F:	M: 0.2,	
		Australia	97		2.9				0.4	1.8	F: 0	
(3)	non-	New	1972-83	> 14	0.55‡	0.43‡	0.15‡	0.23‡	0.26‡	0.42 ‡	0.16‡	0.26‡
	Polynesian	Zealand		years								
	Polynesian				0.22‡	0.58‡	0.74‡	0.7‡	0.35‡	0.3 ‡	0.25‡	0.53‡
(16)	All	French	2007	<15								18
		Polynesia		years								
(17)	Caucasian	East	1988	> 15	4.02							
		France		years	(3.59,							
					4.46)¶							
(19)	Caucasian	Rhone-	1987-88	All	2.7 (2.5,	1.2 (1.0, 1.4)	0.6 (0.5, 0.7)			0.7 (0.6,		
		Alps,			3.0)¶	¶	¶			0.9)¶		
		France										
(18)	Caucasian	Picardy,	1976-80	> 15	0.95	0.52	0.83 (0.50,		0.18	0.24		
		France		years	(0.60,	(0.26,0.74)¶	1.10)¶			(0.07,0.40)		
					1.24)¶					¶		
			1981-85		1.51	1.03	0.26		0.65	0.23		
					(1.10,1.93	(0.69,1.38)¶	(0.09,0.44)¶			(0.00,0.19)		

Table 1A: Incidence rates by type of glomerulonephritis; rates are crude and presented in units of /100 000 /year unlessotherwise specified.

Ref	Race	Location	Time	Age	IgA	Membranous	Membranopr	Mesangial	Minimal	Focal	Anti-	Postinfecti
			period	range	nephropa	nephropathy	oliferative	proliferativ	change	segmental	GBM	ous GN
					thy		GN	e GN	disease	glomerulos	disease	
										clerosis		
)¶					¶		
(6)	Caucasian	Saint	1976-85	20-79	2.8							
		Brieuc,										
		France										
			1986-95	20-79	2.8							
			1996-	≥ 20	2.6							
			2002									
(6)	Caucasian	St. Brieuc,	1976-90	0-79			M: 0.9, F: 0.3					
		France										
(6)	Caucasian	St. Brieuc,	1976-90	10-79	M: 4.8, F:	1.4 (1.1,1.7)¶		0.45		0.8		
		France			1.4, B:			(0.29,0.62)¶		(0.6,1.1)¶		
					2.7							
					(2.2,3.0)¶							
(6)	Caucasian	Saint	1976-85	20-59		1.0						
		Brieuc,										
		France										
				60-79		2.8						
			1986-95	20-59		1.1						
				60-79		3.3						
				00-79		5.5						
			1996-	20-59		0.6						
			2002									
				60-79		1.7						
				≥ 80		0.9						

Fr casian Si Fr	St. Brieuc, France St. Brieuc, France	period 1976-90 1976-90	range > 60 years	nephropa thy 1.0	nephropathy	oliferative GN	proliferativ e GN	change disease	segmental glomerulos	GBM disease	ous GN
Fr casian Si Fr	rance St. Brieuc,					GN	e GN	disease	-	disease	
Fr casian Si Fr	rance St. Brieuc,			1.0							
Fr casian Si Fr	rance St. Brieuc,			1.0					clerosis		
casian SI Fr	St. Brieuc,	1976-90	vears		2.5 (1.7,3.3)¶	0.4 (0.1,0.8)¶		0.7			
Fr	-	1976-90	, caro	(0.5,1.6)¶				(0.3,1.2)¶			
	rance	1970-90	10-79			0.55					0.55
casian Sa	runce										
	Saint	1976-85	20-59			0.9					
В	Brieuc,										
Fr	rance										
			60-79			0.5					
		1986-95	20-59			0.1					
			60-79			0.2					
		1996-	20-59			0.2					
		2002									
casian It	taly	1993	All	0.84	0.49	0.14	0.19	0.16	0.23	0.01	0.07
casian It	taly	1992-94	0-15	0.31	0.015	0.075	0.16	0.23	0.14		0.035
casian Ci	City or	1990-94	≥ 15	M: 2.27,	M: 1.84, F:	M: 0.24, F:		M: 0.36,	M: 0.62, F:		
Pr	rovince			F: 0.67,	0.79, B: 1.31	0.14, B: 0.19		F: 0.27,	0.49, B:		
of	of Turin,			B: 1.47				B: 0.32	0.55		
It	taly										
casian W	Vestern	1980-	All	UH 5.0	UH 1.4						
Fi	inland	2000			CH 0.8						
c	asian I asian I asian C F c I asian V	FranceasianItalyasianItalyasianCity orProvinceof Turin,ItalyItaly	France I986-95 1986-95 1996- 2002 asian Italy I993 asian Italy I1992-94 Province of Turin, Italy I1990-94 Province of Turin, Italy I1990-94 Province	France $60-79$ $60-79$ $60-79$ $1986-95$ $20-59$ $60-79$ $60-79$ $1996 20-59$ 2002 $20-59$ asian Italy 1993 asian Italy $1992-94$ $0-15$ asian City or $1990-94$ 215 Province 0 114 114 114 $1990-94$ 215 114 114 114	$ \begin{array}{c c c c c c c } & France & & & & & & & & & & & & & & & & & & &$	France 60-79 1986-95 20-59 60-79 60-79 1996- 20-59 2002 20-59 asian Italy 1993 asian Italy 1992-94 asian City or 1990-94 Province 1990-94 of Turin, 1990-94 Italy 1990-94 All 0.31 0.79, B: 1.31 of Turin, 1980- Italy 1980- All UH 5.0 UH 1.4	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	France 60-79 60-79 0.5 1986-95 20-59 0.1 0.1 60-79 60-79 0.2 0.2 1996- 20-59 0.2 0.2 2002 2002 0.2 0.2 asian Italy 1993 All 0.84 0.49 0.14 0.19 asian Italy 1992-94 0-15 0.31 0.015 0.075 0.16 asian Italy 1990-94 2-15 M: 2.27, F: 0.67, F: 0.79, B: 1.31 0.14, B: 0.19 0.14, B: 0.19 asian City or 1990-94 2-15 M: 2.27, F: 0.67, B: 1.31 0.14, B: 0.19 0.14, B: 0.19 asian Vestern 1980- All UH 5.0 UH 1.4 UH 5.0	France 60-79 60-79 0.5 1986-95 20-59 0.1 0.1 60-79 0.2 0.2 0.2 1996- 20-59 0.2 0.2 2002 20-59 0.2 0.2 asian Italy 1993 All 0.84 0.49 0.14 0.19 0.16 asian Italy 1992-94 0-15 0.31 0.015 0.075 0.16 0.23 asian City or 1990-94 215 M: 2.27, F: 0.67, of Turin, Italy M: 1.84, F: M: 0.24, F: M: 0.36, 0.14, B: 0.19 M: 0.36, F: 0.27, B: 1.47 asian Western 1980- All UH 5.0 UH 1.4 UH 5.0 UH 1.4	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	France France 60-79 70-79

Ref	Race	Location	Time	Age	IgA	Membranous	Membranopr	Mesangial	Minimal	Focal	Anti-	Postinfecti
			period	range	nephropa	nephropathy	oliferative	proliferativ	change	segmental	GBM	ous GN
					thy		GN	e GN	disease	glomerulos	disease	
										clerosis		
(44)	Caucasian	Denmark	1985-97	All		0.48	0.21	1.08	0.73	0.57		
						(0.4,0.51)¶	(0.16,0.24)¶	(0.94,1.01)¶	(0.62,0.75	(0.48,0.59)¶		
									P(
(21)	Caucasian	Heerlen,	1978-84	16-65	1.9	0.9			0.6	0.9		
		Maastricht		years								
		and										
		Sittard,										
		The										
		Netherlan										
		ds										
(14)	Caucasian	Spain	1994-99	All ages	0.79	0.62	0.36		0.48	0.64		
(20)	Caucasian	Spain	1977-86	> 14	0.35	0.18 (0.17,						
				years	(0.33,0.37	0.20)§						
)§							
(29)	All	Northern	1976-05	>16	0.99	0.75 (0.66,	Type 1: 0.24		0.25 0.20,	0.15 (0.11,		
		Ireland,		years	(0.88,	0.84) ¶	(0.19, 0.29)¶		0.30)¶	0.18) ¶		
		UK			1.09)¶		Type 2: 0.03					
							(0.01, 0.04)¶					
(23)		Leicesters	1973-82	0-15								
Prima		hire, UK										
ry	Asian (i.e.	City							12.1			
nephr	11.9% of	Rest of							6.2			
otic	population	country										
syndr)											
ome	Non-Asian	City							0.4			
		Rest of							1.6			

Ref	Race	Location	Time period	Age range	IgA nephropa thy	Membranous nephropathy	Membranopr oliferative GN	Mesangial proliferativ e GN	Minimal change disease	Focal segmental glomerulos	Anti- GBM disease	Postinfecti ous GN
										clerosis		
		country										
(24)	All	Yorkshire,	1987-98	0-15					2.3 (2.0-			
Prima		UK							2.6)			
ry												
nephr	South								7.4 (5.3-			
otic	Asian								9.5)			
syndr	Not South								1.6 (1.3-			
ome	Asian								1.8)			
(22)	All	Birmingha	1979-83	0-16					5.3 (3.7,			
		m, UK							7.0)¶			
	Afro-								3.4 (0,			
	Caribbean								8.1)¶			
	Asian								15.6 (9.6,			
									21.9)¶			
	Asian								2.6 (1.3,			
									4.0)¶			
(27)	Caucasian	Czech	1994-	All	1.12	0.30 (0.26,	0.15 (0.13,	0.37 (0.32,	0.40	0.35 (0.31,		
		Republic	2000		(1.04,	0.34)¶	0.18)¶	0.40)¶	(0.35,	0.39)¶		
					1.19)¶				0.45)¶			
(26)	Caucasian	Moldova	2004	>18		0.53	0.93	1	0.73	0.33		
		and		years								
		Banat,										
		Romania										
(30)	All	Serbia	1987-06	>18	0.85	1.24	0.61	1.08	0.19	1.11		
				years								

Ref	Race	Location	Time	Age	IgA nephropa	Membranous nephropathy	Membranopr oliferative	Mesangial proliferativ e GN	Minimal	Focal	Anti- GBM disease	Postinfecti ous GN
			period	range					change	segmental		
					thy		GN		disease	glomerulos		
										clerosis		
(38)	All	Bahrain	1990-	All	0.01	0.35 (0.23,	0.38 (0.25,		0.79	0.62 (0.45,		0.08 (0.02,
			2002		(0.00,	0.48)§	0.50)§		(0.60,	0.79) §		0.14)§
					0.03) §				0.97)§			
(39)	Arabian	Bahrain	2003-06	All	0.26	0.22 (0.04,	0.37 (0.14,		0.15	0.30 (0.09,		0.11 (0.00,
	and non-				(0.07,0.46	0.40)§	0.60)§		(0.00,	0.51)§		0.24) §
	Arabian)§				0.30)§			
(13)	Kuwaiti	Kuwait	1995-	All	M: 2.5, F:	M: 0.8, F:1.4,			9.2	M: 1.8, F:		
	national		2001		1.0, B:	B: 1.1				1.1, B: 1.4		
					1.7							
(40)	Kuwaiti	Kuwait	1981-85	0-15					7.2 (5.3,			
	national								9.1)¶			
(37)	Arabian ¹	Benghazi,	1980-82	0-15					11.6 (6.4,			
		Libya							16.9)¶			
(31)	Caucasian	Ottawa-	1985-93	6		0.05 (0, 0.14)	0.05 (0, 0.14)		2.81	0.37 (0.11,		
		Hull		months		¶	¶		(2.10,	0.63)¶		
		region,		- 19					3.52)¶			
		Canada		years								
			1993-			0.09 (0, 0.21)	0		2.47	0.94 (0.54,		
			2002			¶			(1.81,	1.34)¶		
									3.12)¶			
(32)	Caucasian	Central	1975-84	All	M: 0.92,							
		and			F: 0.34,							
		Eastern			B: 0.62							
		Kentucky,			(0.47,0.76							
		US			P(

Ref	Race	Location	Time	Age	IgA	Membranous	Membranopr	Mesangial	Minimal	Focal	Anti-	Postinfecti
			period	range	nephropa	nephropathy	oliferative	proliferativ	change	segmental	GBM	ous GN
					thy		GN	e GN	disease	glomerulos clerosis	disease	
			1985-94		M: 1.43,							
					F: 0.65,							
					B: 1.02							
					(0.84,1.21							
)¶(
(35)	Caucasian	Olmsted	1974-03	All	1.4 (1.0,	0.7 (0.4, 1.0)	0.4 (0.2, 0.7)		0.3 (0.1,	1.1 (0.7,		
		County, US			1.8)				0.5)	1.5)		
(36)	All	New	2000-05	All	0.93							
		Mexico			(0.75.							
					1.11) ¶							
(34)	Caucasian	New	1994-03	1-18					1.81			
Disea	and	Orleans,		years					(1.53,			
se	African	US							2.09) ¶			
define	American											
d as												
nephr												
otic												
syndr												
ome												
(33)	African-	Shelby	1975-84	< 18	0.08							
	American	County,		years	(0.01,							
		Tennessee			0.46)							
		, US										
	Caucasian				0.56 (0.2							
					1.22)							

Ref	Race	Location	Time	Age	IgA	Membranous	Membranopr	Mesangial	Minimal	Focal	Anti-	Postinfecti
			period	range	nephropa	nephropathy	oliferative	proliferativ	change	segmental	GBM	ous GN
					thy		GN	e GN	disease	glomerulos	disease	
										clerosis		
	African-		1985-94		0.57							
	American				(0.23,							
					1.18)							
	Caucasian				0.3 (0.06,							
					0.87)							
(41)	All	Venezuela	1998	<15	0.03 (0,				2.4 (2.0,			
				years	0.1)§¶				2.7) §¶			
(15)	All	Uruguay	1990-94	All	0.24	0.16	0.06		0.4	0.67		
			1995-99		0.46	0.31	0.10		0.55	0.99		
					0.45	0.40	0.14		0.46	0.64		

Notes: *‡* incidence rate adjusted for age; § incidence rate calculated from data given in the paper; ¶ CI calculated from data given in paper using standard Normal distribution; ¹ Terminology as given in paper; UH University hospital; CH central hospital