Increasing prevalence of Addison’s disease in German females: health insurance data 2008–2012

Gesine Meyer, Kathrin Neumann1, Klaus Badenhoop and Roland Linder1

Division of Endocrinology, Department of Medicine 1, University Hospital, Goethe-University Frankfurt, Theodor-Stern-Kai 7, D-60590 Frankfurt, Germany and 1WINEG Scientific Institute of the TK for Benefit and Efficiency in Health Care, Hamburg, Germany

Abstract

Objective: Our objective was to investigate the epidemiology of autoimmune Addison’s disease (AD) in Germany.

Design: Routine data were analyzed from the Statutory Health Insurance (SHI) database of the Techniker Krankenkasse (TK) for an observation period from 01/01/2008 to 31/12/2012. The TK is one of the largest German health care insurance providers covering more than 10% of the German population.

Subjects and methods: Between 2008 and 2012, a total of 2477 diagnoses of primary adrenal failure were recorded in the SHI database. After exclusion of secondary, iatrogenic or other non-idiopathic forms and after adjustment for incomplete data sets, 1364 diagnoses of autoimmune-mediated AD remained.

Results: The prevalence of AD in our cohort showed a steady increase from 82 per million in 2008 to 87 per million in 2012. On average, the prevalence rose about 1.8% per year, and due to a pronounced increase (2.7%) in females. The prevalence was lower in men (63–68 per million) than in women (96–108 per million). Autoimmune comorbidities were found in 46.5% of AD patients. Adrenal crises were documented with a frequency of 14–17/100 patient years.

Conclusions: These data provide a first epidemiological profile of this rare and perilous endocrine disease in Germany. Although the prevalence of AD appears lower than in the Scandinavian countries, the increasing figures in females over the last 5 years warrant further investigations. Furthermore, adrenal crises pose a considerable burden. Hereby, we can show that health insurance data provide a valuable tool for epidemiological studies in the absence of national registries.

Introduction

Addison’s disease (AD) is a chronic disease leading to a deficient production of glucocorticoids, mineralocorticoids, and androgens in the adrenal cortex. The disease is rare and valid epidemiological data are difficult to collect. For the European countries, the prevalence is estimated at 93–144 cases per million (1, 2). These estimations, however, are based on few data from rather small populations – less than one million – in single European regions (3, 4, 5, 6). Most convincing data come from the Scandinavian countries, where national patient registries exist (7, 8). Norway and Sweden, moreover, have the highest prevalence for autoimmune AD with 144 and 136 per million respectively (6, 7, 8). Primary adrenal insufficiency occurs more frequently in women compared with men with an odds ratio around 1.2 for women (1, 5).

The incidence of AD in Europe is estimated at 4.4–6.2 per million population per year. Several recent studies have pointed to an increasing incidence of the disease in the last decades (1, 2, 4, 6). Very recently, a Swedish study, based on the Swedish National Prescribed Drug Register, has shown a distinct increase in yearly prevalence for autoimmune AD from 12.2 to 13.1/100 000 person years in the period 2005–2009 (8). Considering the continuous decline in tuberculous adrenalitis, one of the major causes
Subjects and methods

The SHI database of the TK provides a documentation of diagnoses by all members with a doctor contact, comprising data of inpatient and ambulatory medical treatment as well as outpatient surgery.

Diagnoses are encoded by the International Statistical Classification of Diseases and Related Health Problems (ICD-10). These data were evaluated by the Scientific Institute Wissenschaftliches Institut der TK für Nutzen und Effizienz im Gesundheitswesen – WINEG.

We analyzed routine data from the TK. ICD-codes as accounting data are transferred in the SHI database automatically. As all data are linked to the patients’ insurance registration number, they are individual related and not exchangeable. Altogether, in the observation period from 01/01/2008 to 31/12/2012, more than 30 million men were evaluated each year.

A query for ICD-10 codes E27.1 (primary adrenal failure), E27.2 (adrenal crisis), and E31.0 (APS) was implemented, the latter by filtering for the medications containing hydrocortisone (ATC code H02AB09) and/or fludrocortisone (ATC code H02AA02) in the year of diagnosis. Subsequently, all causes of secondary, iatrogenic, infectious, traumatic, or other non-idiopathic forms of primary adrenal failure were excluded by filter out ICD-10 codes E23.0 (pituitary failure), E22 (pituitary hyperfunction), E24 (Cushing’s syndrome), D35.2, D35.3, D44.3, and D44.4 (pituitary tumors), E27.2 (primary adrenal failure), E89.6 (adrenal failure caused by medical measure), A18.7 (tuberculous adrenalitis), A39.1 (Waterhouse–Friderichsen syndrome), E27.8 (miscellaneous disease of adrenal gland), E27.4 (atrophic adrenitis), E29 (benign adrenal tumor), C79.7 (adrenal metastasis), and C74.9 (malignant adrenal tumor).

Patients with AD, defined in this manner as autoimmune-mediated AD, were screened for associations with other autoimmune diseases, searching for codes K29.4 (atrophic gastritis), K75.4 (autoimmune hepatitis), K90 (celiac disease), M35 (collagen disease), E05 (Graves’ disease), E06 (Hashimoto’s thyroiditis), E20 (hypoparathyroidism), M32.1/.8/.9 (lupus erythematosus), D51.0 (pernicious anemia), E28.3 (primary ovarian failure), M05/M06 (rheumatoid arthritis), E10 (type 1 diabetes mellitus), and L80 (vitiligo).
Data were collected annually for the years 2008–2012. All data were adjusted to the general population of Germany for age and sex, using the data of the German Federal Office of Statistics from 2010.

In an analogous manner, we implemented a query for the autoimmune diseases, type 1 diabetes mellitus (ICD-10 code E10) and vitiligo (ICD-10 code L80), in all TK-insurants. These data were collected annually for the years 2009–2011. The study was approved by the Local Ethical Committee.

Results

Between 2008 and 2012, 2477 diagnoses of primary adrenal failure were recorded in the SHI database. After exclusion of secondary, iatrogenic, or other non-idiopathic forms and after adjustment for incomplete data sets, 1364 diagnoses of autoimmune-mediated AD remained.

Therefore, in our cohort, representing ~10% of the German population, the prevalence of autoimmune-mediated AD was found to range between 80 and 87 per million. The prevalence showed an increase from 82 per million in 2008 to 87 per million in 2012. On average, the prevalence rose about 1.8% per year (0.2% in men and 2.7% in women; Fig. 1).

Diagnosis was documented in n=536 male and n=828 female insurants. The percentage of affected males varied between 38.2 and 41.5%. Altogether, the prevalence of autoimmune AD was lower in men (63–68 per million) than in women (96–108 per million) with an odds ratio of 1.6 (95% CI: 1.3–1.8) for women. Male patients were 5–94 years old (mean age: 51 years) compared with female patients aged 4–95 years (mean age: 48 years). Adrenal crises were documented with a frequency of 14–17/100 patient years.

Autoimmune comorbidities were found in 46.5% (n=634). Commonest were autoimmune thyroid diseases and type 1 diabetes mellitus, followed by rheumatoid diseases, vitiligo, atrophic gastritis, and primary ovarian failure (Fig. 2). Individual patients had up to five additional autoimmune diseases (+1, 61.4%; +2, 29.1%; +3, 7.3%; +4, 1.9%; and +5, 0.3%).

Compared with TK-insurants without AD, patients with AD were found to have a severe risk for diverse additional autoimmune-mediated diseases of endocrine and non-endocrine organs up to an odds ratio of >30. Hereby, autoimmune thyroid diseases not only showed the highest risk, diagnosed in 44.5%, but also type 1 diabetes was found in nearly 12% of AD patients. Rare conditions like pernicious anemia, autoimmune hypoparathyroidism, and autoimmune hepatitis illustrate the broad spectrum of autoimmunity in AD patients (Table 1).

To compare potential changes in the prevalence of other autoimmune diseases, we specifically investigated the change in the prevalence of type 1 diabetes mellitus: its prevalence in all TK-insurants lay between 7961 and 8313 per million and did not change significantly over the years 2009–2011. A similar result was seen for vitiligo with a prevalence between 1395 and 1422 per million, with no significant change.

Discussion

Epidemiological data in AD have been provided not only by either Scandinavian registries but also through investigator-initiated databases in the academic centers with a large referral area (3, 4, 5). We provide such data for the first time through a SHI. Such a database implies potential limitations. As coding is the basis for reimbursement both in hospitals and in outpatient care, we assume a correct code for rare AD both by primary care physicians as well as by specialists, such as in endocrinology. However, incomplete diagnosis coding cannot be ruled out particularly in autoimmune polyendocrine symptom (APS) patients. In this study, we filtered out other forms of adrenal failure to minimize this error rate and included only those APS patients with an AD steroid substitution profile. Furthermore, we confined the query for differential diagnoses to the year quarter, in which the diagnosis
of AD was encoded. This approach bears a remaining risk for incorrect coding particularly missing some AD patients. We therefore assume a conservative estimation of the true prevalence.

In the SHI database of the TK, patients, and therefore their diagnoses, are recorded only when they had an inpatient or outpatient doctor contact due to this diagnosis in a particular year. For this reason, our results reveal indications for the prevalence of autoimmune AD related to a distinct year and not for the incidence of this disease. As patients with primary adrenal insufficiency depend on a lifelong corticosteroid supplementation therapy and this medication is available only on prescription in Germany, each Addison patient has at least one medical contact per year.

The retrieval system using data from a large statutory health care insurance was valid and comparable to those from other countries with national diagnostic ascertainment such as Sweden and Norway (6, 7, 8). Thereby this novel tool allows epidemiological studies on rare diseases in the absence of national registries. Moreover, health care providers would benefit by allocating medical resources based on reliable diagnostic data and can improve the long-term management of both rare and more common chronic disorders.

We found a prevalence of autoimmune AD in Germany between 82 and 87 per million. This prevalence appears lower than in the Scandinavian countries, Norway (144 per million) and Sweden (136 per million) (6, 7). It is also lower than most of the estimations for other European countries so far (3, 4, 5, 6). As the latter figures are based on few data from very small populations, their validity is limited. Owing to the large size of the TK-insurants, covering nearly 10% of the German population, and the

Table 1  Odds ratio (OR) for 13 autoimmune comorbidities in individuals with autoimmune Addison’s disease (AD) compared with individuals without AD (controls).

<table>
<thead>
<tr>
<th>ICD</th>
<th>Individuals with AD in 2012 (n=712)</th>
<th>Controls (n=8 540 805) in 2012</th>
<th>OR</th>
<th>Lower 95% CI</th>
<th>Upper 95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>With ICD</td>
<td>Without ICD</td>
<td>With ICD</td>
<td>Without ICD</td>
<td></td>
</tr>
<tr>
<td>E06.3: Hashimoto’s thyroiditis</td>
<td>268</td>
<td>444</td>
<td>171 663</td>
<td>8 369 142</td>
<td>29.4</td>
</tr>
<tr>
<td>E10: type 1 diabetes mellitus</td>
<td>85</td>
<td>627</td>
<td>74 506</td>
<td>8 466 299</td>
<td>15.4</td>
</tr>
<tr>
<td>M05/M06: rheumatoid arthritis</td>
<td>61</td>
<td>651</td>
<td>86 372</td>
<td>8 454 433</td>
<td>9.2</td>
</tr>
<tr>
<td>E05.0: Graves’ disease</td>
<td>49</td>
<td>663</td>
<td>37 003</td>
<td>8 503 802</td>
<td>17.0</td>
</tr>
<tr>
<td>M35.0: collagen disease</td>
<td>29</td>
<td>683</td>
<td>81 926</td>
<td>8 458 879</td>
<td>4.4</td>
</tr>
<tr>
<td>E28.3: primary ovarian failure</td>
<td>13</td>
<td>699</td>
<td>21 944</td>
<td>8 518 861</td>
<td>7.2</td>
</tr>
<tr>
<td>L80: vitiligo</td>
<td>21</td>
<td>691</td>
<td>14 488</td>
<td>8 526 317</td>
<td>17.9</td>
</tr>
<tr>
<td>D51.0: pernicious anemia</td>
<td>18</td>
<td>694</td>
<td>6773</td>
<td>8 534 032</td>
<td>32.7</td>
</tr>
<tr>
<td>K29.4: atrophic gastritis</td>
<td>14</td>
<td>698</td>
<td>6919</td>
<td>8 533 886</td>
<td>24.7</td>
</tr>
<tr>
<td>K90.0: celiac disease</td>
<td>14</td>
<td>698</td>
<td>11 730</td>
<td>8 529 075</td>
<td>14.6</td>
</tr>
<tr>
<td>K75.4: autoimmune hepatitis</td>
<td>7</td>
<td>705</td>
<td>2453</td>
<td>8 538 352</td>
<td>34.6</td>
</tr>
<tr>
<td>E20: primary hypoparathyroidism</td>
<td>7</td>
<td>705</td>
<td>3094</td>
<td>8 537 711</td>
<td>27.4</td>
</tr>
<tr>
<td>M32.1/M32.8/M32.9: lupus erythematosus</td>
<td>5</td>
<td>707</td>
<td>4090</td>
<td>8 536 715</td>
<td>14.8</td>
</tr>
</tbody>
</table>
quantity of analyzed data, we assume that our findings are sufficiently valid. Our data provide a first epidemiological profile of rare and perilous AD in Germany.

In our cohort, adrenal crises were documented with a frequency of 14–17/100 patient-years. This finding exceeds previous observations considerably. Former data of a German population showed about 6.3-crises/100 patient-years calculated by patient questionnaires in a mixed cohort of patients with primary and secondary adrenal insufficiency (10). In agreement with existing data, AD affects more frequently women than men in our cohort with an odds ratio slightly higher than the reported one (1.6 vs 1.2).

Furthermore, we observed an annual increase in prevalence of about 1.8% per year on average. A relevant change in the medical profession’s awareness over the years 2008–2012 for AD as reason for this observation is not apparent. Actually, the awareness is still insufficient. Two recent studies have shown a delay of diagnosis (11, 12). As medical treatment of AD has not changed over decades, we assume that there is no change in life expectancy beyond that of the general population.

Remarkably, the increment of prevalence was nearly exclusively seen in female individuals (2.7% per year in average vs 0.1% in males). To the best of our knowledge, this is the first study of AD that showed a difference in the increasing prevalence for women and men.

An increasing incidence has been observed in AD (1, 2, 4, 6, 8) and several other autoimmune-mediated disorders in recent years, including celiac disease (13), type 1 diabetes (14, 15, 16, 17, 18), and multiple sclerosis (19). Diverse genetic variations predispose to autoimmune AD (20, 21, 22). Most of them are susceptibility factors for autoimmunity in general. However, genetic susceptibility variants are rather frequent in the general population, and their role for an increase in prevalence of autoimmune diseases is probably not relevant because their frequencies have not changed. More likely is the role of environmental factors to cause and/or trigger organ-specific autoimmunity. In recent years, vitamin D deficiency has been implicated as a potential environmental factor for autoimmune disorders (23, 24). We have no data about a possible change in prevalence of vitamin D deficiency in AD patients during the study period, but in patients with type 1 diabetes we do observe such a trend (25). Further investigations in patients with AD are necessary.

Other environmental agents, such as viral infections, cigarette smoking, and pollutants, as well as a reduced microbial burden during early life, have been found to influence the development of autoimmune diseases, presumably by epigenetic changes (26, 27, 28).

A striking finding is that particularly females show an increasing prevalence over the last 5 years. One possible explanation is a growing trend for autoimmunity in general. However, in contrast to preceding studies (14, 15, 16, 17, 18), in our cohort the prevalence data for the much more frequent autoimmune diseases type 1 diabetes mellitus and vitiligo did not change relevantly over the years 2009–2011, but the triggering and/or causative agents in rare AD may differ from those more common disorders.

Possibly, so far undetected environmental factors may underlie the distinct rise of autoimmune AD in the recent years. The stronger increase in prevalence in women raises the question, which environmental factors may affect females more than males. A stronger increase in type 1 diabetes has recently been described in Sardinian girls who might be more vulnerable to environmental factors (13).

In patients with autoimmune-mediated AD, autoimmune comorbidity is rather frequent and occurs in ~60% of patients. In our cohort, the rate was lower with an autoimmune comorbidity of 46.5%. Conceivably, focusing on the principal diagnosis of AD, comorbidities may not have been coded for in every case. Furthermore, analysis of data was limited to the years 2008–2012, so future manifestations of autoimmune-mediated diseases in patients with AD could not be regarded. This would lead to an underestimation especially of autoimmune comorbidity, which is possibly more frequent than we observe.

We found a severe risk for diverse additional autoimmune-mediated diseases of endocrine and nonendocrine organs in patients with autoimmune AD up to an odds ratio of >30 compared with individuals without AD. This impact was expectedly strong for frequent autoimmune diseases like Hashimoto’s thyroiditis (odds ratio 29.4), but notably also for rare disorders like pernicious anemia (odds ratio 32.7), autoimmune hepatitis (odds ratio 34.6), and atrophic gastritis (odd ratio 24.7).

Our cohort is confined to German health care insurants. Further research in other countries is necessary to prove the increasing prevalence to be a global phenomenon. However, in order to compare epidemiological data for rare autoimmune AD, a standardized procedure of operationalization is required. With this study, we provide a differentiated approach for standardization of inclusion as well as exclusion criteria.
Our findings emphasize the importance of a heightened clinical awareness not only to correctly diagnose the growing number of AD patients but also to recognize their comorbidities, especially autoimmune thyroid diseases, and also rare autoimmune diseases. Furthermore, we identify a relevant risk of adrenal crises with a quantity of about 14–17/100 patient years. As every adrenal crisis is potentially life threatening and harbors the risk of recurrence, this result warrants intensified measures for prevention.

Despite the rising prevalence, the disease remains rare (<2/5000) and thereby the medical professions need to be alerted that these patients require timely and adequate treatment and follow-up. Furthermore, elucidating the triggering factors for these epidemiological changes will lead to a better understanding of its pathophysiology and possibly new roads to immune therapy.

Declaration of interest
G Meyer, K Neumann, and K Badenhoop have nothing to declare. R Linder is employed by WINEG Scientific Institute of the Techniker Krankenkasse for Benefit and Efficiency in Health Care, which investigates the value of innovations and new programatic approaches within the Statutory Health Insurance framework.

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