Anticholinergic drug burden according to the anticholinergic drug scale and the German anticholinergic burden and their impact on cognitive function in multimorbid elderly German people: a multicentre observational study


ABSTRACT

Objectives The aims of our study were to examine the anticholinergic drug use and to assess the association between anticholinergic burden and cognitive function in the multimorbid elderly patients of the MultiCare cohort.

Setting MultiCare was conducted as a longitudinal cohort study in primary care, located in eight different study centres in Germany.

Participants 3189 patients (59.3% female).

Primary and secondary outcome measures Baseline data were used for the following analyses. Drugs were classified according to the well-established anticholinergic drug scale (ADS) and the recently published German anticholinergic burden (German ACB). Cognitive function was measured using a letter digit substitution test (LDST) and a mixed-effect multivariate linear regression was performed to calculate the influence of anticholinergic burden on the cognitive function.

Results Patients used 1764 anticholinergic drugs according to ADS and 2750 anticholnergics according to the German ACB score (prevalence 38.4% and 53.7%, respectively). The mean ADS score was 0.8 (±1.3), and the mean German ACB score was 1.2 (±1.6) per patient. The most common ADS anticholinergic was furosemide (5.8%) and the most common ACB anticholinergic was metformin (13.7%). The majority of the identified anticholinergics were drugs with low anticholinergic potential: 80.2% (ADS) and 73.4% (ACB), respectively. An increasing ADS and German ACB score was associated with reduced cognitive function according to ADS and German ACB score. We especially need to gain greater awareness for the contribution of drugs with low anticholinergic potential from the cardiovascular system. As anticholinergic drug use is associated with reduced cognitive function in multimorbid elderly patients, the importance of rational prescribing and also deprescribing needs to be further evaluated.

Trial registration number ISRCTN89818205.

INTRODUCTION

The greater number of people in the population surviving until very late life leads to a challenge to the provision of healthcare, particularly given the proportion of older people that live with multiple comorbidities. These in turn often lead to polypharmacy, which is commonly defined as the coapplication or coprescription of five or more drugs at the same time. Apart from this, it is also known that multimorbid elderly patients are at a higher risk for taking anticholinergic drugs or drugs that have anticholinergic side effects. Besides classic anticholinergic substances—for example, drugs for urinary incontinence, chronic obstructive pulmonary disease or Morbus Parkinson—a lot of drugs...
lead to anticholinergic adverse drug reactions (ADRs). These ADR and also the intended anticholinergic effects are evoked by the binding of drugs to one of the five muscarinic receptors in autonomous nervous system and especially blocking the parasympathetic nervous system. Particularly elderly people are more vulnerable towards anticholinergic ADR because of an age-related decreased cholinergic transmission and a poorer metabolism and/or elimination of those substances. Therefore, there is some evidence that the use of anticholinergic drugs or drugs with anticholinergic activity is associated with a higher risk of falls, hospitalisation and even mortality in elderly patients. Anticholinergic drug use is also associated with cognitive impairments and dementia. Moreover, the use of anticholinergics leads to less self-dependency and a decrease in functional status. Likewise, patients might suffer from typical anticholinergic side effects as mental confusion, tremor, visual disturbances, delirium, dry mouth and urinary retention.

The anticholinergic burden, the cumulative effect of using multiple drugs with anticholinergic activity simultaneously, can be calculated with the help of different lists. In the most common lists, drugs are categorised in none, low, moderate or high anticholinergic activity (zero to three points). The gained scores are summed up, and when the score is greater or equal three points, one should consider to use alternative drugs or a dose reduction. Some lists additionally include the daily dose. The number of included drugs varies between the scores and the scoring bases on different methods, for example, with regard to the drug’s potency and efficiency or to its exposure. With regard to the association of anticholinergic burdens on the cognitive function in elderly people, conflicting results have been published. As the different published tools rate drugs quite differently and on different bases, we decided to use two different tools to evaluate anticholinergic burden of our patient collective. The anticholinergic drug scale (ADS) developed by Carnahan et al is validated against serum anticholinergic activity (SAA), and high SAA levels are associated with cognitive impairments. Furthermore, the ADS score is a well-established tool to identify drugs with anticholinergic activity. Kiesel et al developed the German anticholinergic burden score (German ACB) especially for the German drug market in order to improve the routine prescribing in geriatric patients for the German population. To the best of our knowledge, there is no study that compares the ADS score with the German ACB score to investigate the anticholinergic burden of elderly multimorbid patients and pointing out the effect of anticholinergic drug use on the cognitive function. As far as we know, there is still limited data about the influence of anticholinergic drugs on the cognitive function from large European patient cohorts.

The German MultiCare cohort offers ideal conditions, as the study was conducted in order to examine the influence and effects of multimorbidity in multimorbid elderly patients in primary care. Patients and general practitioners (GPs) were interviewed about morbidities, prescribed and over-the-counter (OTC) medication, socioeconomic status, risk factors, health status and functional status, among others.

The aims of our study were: (1) to identify anticholinergics and drugs with anticholinergic activity with the ADS and the German ACB score (2) and to show the effect of the anticholinergic burden measured with the German ACB score on the cognitive function and compare those findings with the well-established ADS score.

METHODS

Study design

The MultiCare study was carried out as a multicentre, observational cohort study in primary care. Baseline data collection started in July 2008, and three follow-ups were performed, and each recruitment wave took 15 months. For our analysis, the baseline assessment of 3189 patients, collected between 21 July 2008 and 6 November 2009, was used. The recruitment took place in eight study centres in Germany (Bonn, Duesseldorf, Frankfurt/Main, Hamburg, Jena, Leipzig, Mannheim and Munich). Multimorbid patients were randomly selected from the GPs’ electronic files in 158 practices. Patients were included if they had at least three diagnosed chronic diseases and were between 65 and 85 years old. Exclusion criteria were: (1) nursing home patients, (2) blindness, (3) deafness, (4) missing capacity to consent, particularly patients with dementia, (5) all patients who had an expected life expectancy of less than 3 months, (6) insufficient ability to read and speak German, (7) patients who participate in other studies and (8) patients poorly known by the physician. A total of 7172 patients out of 50 786 patients from the GPs were contacted for informed consent after screening for inclusion and exclusion criteria. With a total response rate of 46.2%, 3317 patients were included, and after excluding 128 patients because they died before the baseline interview or due to other reasons, 3189 patients remained in the cohort. Standardised interviews and tests, at patients home, were conducted with the remaining 3189 patients to collect data about sex, age, education, income and cognitive skills by using the letter digit substitution test (LDST). Additionally, a brown bag review—capturing all prescription and OTC drugs the patients used on a regular basis or on demand—was performed to collect information about patients’ medication. Information about morbidity was gained with the help of standardised GP interviews. Schäfer et al previously published detailed information on the exact study design (online supplemental file 1).

Anticholinergic burden classification, descriptive results and subgroup analysis

Prescription and OTC drugs were gathered via brown bag review, and the drugs were classified analogous to the anatomical therapeutic classification (ATC) system.
We used Excel 2016 (Microsoft Office 2016, Redmond, USA) to rate the anticholinergic drugs according to the German ACB and ADS scores.

The German ACB score was especially developed for the German drug market by Kiesel et al. Drugs were classified as drugs with anticholinergic activity with the help of a systematic literature research in PubMed and a subsequently evaluation by experts. The German ACB score comprises 507 substances, whereby 356 drugs have no anticholinergic effect (ACB score=0), 104 drugs are scored as weak (ACB score=1), 18 drugs are scored as moderate (ACB score=2) and 29 drugs are scored as having strong (ACB score=3) anticholinergic effects.

The ADS comprises 413 substances and is based on expert opinions. The ADS score categorises drugs into four different levels. Level 0 with no anticholinergic effect (296 substances), 71 level 1 drugs with a weak anticholinergic effect, 12 level 2 drugs with a moderate anticholinergic effect and 34 level 3 drugs with a strong anticholinergic effect.

The anticholinergic burden was calculated by summing up the individual anticholinergic scores of each patient, according to both anticholinergic scores individually.

For gaining the results for the subgroup analysis, a t-test with STATA V.15.1 (StataCorp, College Station, USA) was performed. We defined an alpha-level of 5% (p≤0.05) as statistically significant.

**Fit for the Aged (FORTA) classification**

For the classification according to FORTA PIM list, drugs were analysed indication based with QlikView 11.20 (QlikTech, Radnor, USA). The FORTA list comprises 296 drugs used in the treatment of 30 diagnoses or indications. FORTA rated drugs indication based as: A (absolute), B (beneficial), C (careful) and D (don’t). Drugs were classified as a potentially inadequate medication (PIM) when they are a FORTA C or D drug. For FORTA list is used to reveal whether an additional use of an anticholinergic burden classification is necessary or not.

**Association of anticholinergic drug use with the cognitive function**

We performed a multivariate mixed-effect linear regression to calculate the influence of anticholinergic burden detected by the German ACB and the ADS score on the cognitive skills of the patients. Whereby the LDST, as a speed-depending cognitive task, was used to calculate the cognitive skills of the multimorbid elderly patients.

In LDST, patients have to replace letters by numbers in a specific time to show their ability of processing speed, which is an important cognitive ability and expresses normal cognitive development. Sex, age, number of diseases weighted by severity, highest education degree in three groups according to the international CASMIN (comparative analysis of social mobility in industrial nations) classification and household net adjusted disposable income as independent variables into the model were included.

We adjusted the multilevel linear regression for random effects on the study centre and GP practice in order to minimise the regional effect of prescribing because of the eight different study centres (Bonn, Dusseldorf, Frankfurt/Main, Hamburg, Jena, Leipzig, Mannheim and Munich) and the 158 general practices.

The missing values—in LDST (missing values: 243), number of dosages weighted by severity (152), education standard (3) and the income data sets (258)—were imputed via hot deck imputation. The hot deck imputation has been described elsewhere. All analyses were performed with the imputed data sets, and an alpha level of 5% (p≤0.05) was defined as statistically significant. We conducted all statistical test with STATA V.15.1.

**Patient and public involvement**

No patients were involved in setting the research question or the outcome measures, nor were they involved in developing plans for design or implementation of the study. No patients were asked to advise on interpretation or writing up of results. There are no plans to disseminate the results of the research to study participants or the relevant patient community.

**RESULTS**

**Characterisation of the multimorbid elderly patient collective**

A total of 3189 patients aged 65–85 years were included in the study. The mean age was 74.4 (±5.2) years and 59.3% of the patients were female. In total, 24535 drugs including OTC were identified and related to an ATC code. In mean patients used 7.7 (±3.9) drugs (median 7 drugs, range 0–29 drugs). Table 1 summarises the main findings for the ADS and German ACB score. With ADS score, 1764 drugs were identified as anticholinergic for the MultiCare cohort, with a prevalence of anticholinergic drug use of 38.4% (1226). The mean ADS score is 0.8 (±1.3) and 10.5% (334) of all patients had an ADS score of 3 or higher. For ACB, we detected 2750 anticholinergics in total, and the prevalence of anticholinergic drug use is 53.7% (1714). The mean ACB score per patient is 1.2 (±1.6), and 18.1% (567) of all patients had an ACB score of 3 or higher.

As the most common ADS drug, we detected furosemide 5.8% (185) as anticholinergic with low potential. Amitriptyline was identified as the most common anticholinergic ADS drug with a high anticholinergic potential, with 2.8% (88). For the ACB score, we identified metformin with 13.7% (436) as an anticholinergic with low potential, as the most reported ACB anticholinergic in the MultiCare cohort. Tramadol with 3.3% (105) and amitriptyline with 2.8% (88) are the most common ACB anticholinergic drugs with a moderate and high anticholinergic potential, respectively. 80.2% of the anticholinergics according to ADS score and 73.4% of the detected anticholinergics according to ACB score are low potential anticholinergic drugs. ADS score most frequently detected drugs from the cardiovascular system...
(ATC C) with 36.6% (646 drugs, 11 different substances), followed by drugs from the nervous system (ATC N) with 31.9% (563 drugs, 35 different substances). In contrast, drugs from the nervous system make up the largest group of identified ACB anticholinergics, with 50 different substances and 29.5% (812 drugs) in total, followed by the cardiovascular system, with 13 different substances and 25.7% (709 drugs) in total. Considering the distribution of all used drugs within the MultiCare cohort, the proportions of anticholinergic drugs according to ADS and German ACB with regard to drugs from the cardiovascular system are 7.0% and 7.7% and with regard to drugs from the central nervous system are 22.5% and 32.4%, respectively.

In table 2, the top 10 ADS and German ACB anticholinergics and their occurrence in the FORTA PIM list are captured. Two of the top 10 drugs (tiotropium and ipratropium as inhalatives) are not listed in the FORTA list, while only four drugs are classified into the categories C or D.

### Subgroup analysis: age, sex and polypharmacy

Tables 3 and 4 summarise the most important results of the subgroup analysis for both scores. Female patients had a significant higher ADS and German ACB score than male patients (ADS: female: 0.82±1.34 male: 0.65±1.15; p<0.001; ACB: female: 1.30±1.73 male: 1.04±1.42; p<0.001). Patients 80 years old and older had a significant higher ADS score than the patients that are 65 up to 79 years old (p=0.001). In contrast with that, there was no significant effect on the ACB score observed between the two age groups. However, patients using eight drugs or more at the same time had a significant higher ADS and ACB score than patients using less drugs (p<0.001).

### Association of anticholinergic drug use with the cognitive function

On average, patients achieved a mean LDST score of 23 (±7.1) with a range of 0–50 in the LDST, while 51.9% of the patients gained a score between 20 and 29. Figure 1 shows the kernel density estimator of the baseline results of patients LDST, showing the proportion of patients in each category.

## Table 1

<table>
<thead>
<tr>
<th>Anticholinergic drugs per patient</th>
<th>ADS</th>
<th>ACB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of anticholinergic drugs</td>
<td>1764 (7.2%)</td>
<td>2750 (11.2%)</td>
</tr>
<tr>
<td>Prevalence of anticholinergic drug use</td>
<td>38.4%</td>
<td>53.7%</td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>0.6 (±0.9)</td>
<td>0.9 (±1.0)</td>
</tr>
<tr>
<td>Median (range)</td>
<td>0 (0–6)</td>
<td>1 (0–7)</td>
</tr>
<tr>
<td>0 AC per patient</td>
<td>1963 (61.6%)</td>
<td>1475 (46.3%)</td>
</tr>
<tr>
<td>1 AC per patient</td>
<td>846 (26.5%)</td>
<td>1033 (32.4%)</td>
</tr>
<tr>
<td>2 AC per patient</td>
<td>265 (8.3%)</td>
<td>435 (13.6%)</td>
</tr>
<tr>
<td>3 AC per patient</td>
<td>81 (2.5%)</td>
<td>172 (5.4%)</td>
</tr>
<tr>
<td>4 AC per patient</td>
<td>26 (0.8%)</td>
<td>45 (1.4%)</td>
</tr>
<tr>
<td>5 AC per patient</td>
<td>7 (0.2%)</td>
<td>24 (0.8%)</td>
</tr>
<tr>
<td>6 AC per patient</td>
<td>1 (0.03%)</td>
<td>4 (0.1%)</td>
</tr>
<tr>
<td>7 AC per patient</td>
<td>–</td>
<td>1 (0.03%)</td>
</tr>
</tbody>
</table>

## Table 2

<table>
<thead>
<tr>
<th>Anticholinergic score per patient</th>
<th>ADS</th>
<th>ACB</th>
<th>FORTA PIM (categories)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (SD)</td>
<td>0.8 (±1.3)</td>
<td>1.2 (±1.6)</td>
<td></td>
</tr>
<tr>
<td>Median (range)</td>
<td>0 (0–11)</td>
<td>1 (0–11)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 0</td>
<td>1963 (61.6%)</td>
<td>1475 (46.3%)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 1</td>
<td>682 (21.4%)</td>
<td>802 (25.1)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 2</td>
<td>210 (6.6%)</td>
<td>345 (10.8%)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 3</td>
<td>179 (5.6%)</td>
<td>272 (8.5%)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 4</td>
<td>86 (2.7%)</td>
<td>140 (4.4%)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 5</td>
<td>36 (1.1%)</td>
<td>67 (2.1%)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 6</td>
<td>23 (0.7%)</td>
<td>45 (1.4%)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 7</td>
<td>5 (0.2%)</td>
<td>21 (0.7%)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 8</td>
<td>2 (0.1%)</td>
<td>14 (0.4%)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 9</td>
<td>2 (0.1%)</td>
<td>5 (0.2%)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 10</td>
<td>–</td>
<td>1 (0.03%)</td>
<td></td>
</tr>
<tr>
<td>Score per patient: 11</td>
<td>1 (0.03%)</td>
<td>2 (0.1%)</td>
<td></td>
</tr>
</tbody>
</table>

AC, anticholinergic.
We evolved two models to express the influence of anticholinergics on the LDST results. Tables 5 and 6 show the outcomes of the multivariate mixed-effect linear regression for the ADS score and the German ACB score. In the first model, not including FORTA PIM, we detected that with increasing ADS score, the ability to complete the LDST decreases significantly with a regression coefficient of \(-0.37\) (p<0.001). Also, the German ACB score could exhibit the effect of worse LDST results with increasing ACB score with a regression coefficient of \(-0.33\) (p<0.001). According to a sensitivity analysis, we added FORTA PIM as a cofounder to the regression model (ADS score: p=0.257, regression coefficient: \(-0.04\); ACB score: p=0.518; regression coefficient: \(-0.02\)). By adding FORTA PIM into the second model, the regression coefficient dropped but was still significant: for ADS score, we measured a regression coefficient of \(-0.26\) (p=0.008) and for the German ACB, a score of \(-0.24\) (p=0.003).

### DISCUSSION

#### Statement of principal findings

Our study demonstrates that a huge proportion of multimorbid elderly patients are exposed to anticholinergic drugs or drugs with anticholinergic activity and are consequently affected by the risk of anticholinergic adverse reactions that are associated with decreased cognitive function determined by LDST.

#### Anticholinergic burden classification and risk factors for anticholinergic drug use

In terms of the ADS score, our findings are in good accordance with the literature.29–31 As there is no publication analysing medication with the German ACB score yet, we compared our findings with the gained results of the ADS score and other well-established anticholinergic scores. The results for the German ACB score (mean anticholinergic burden: 1.2±1.6; prevalence: 53.7%) are comparable with our findings with the ADS score (0.8±1.3; 38.4%) and also other anticholinergic scores described in literature (0.5±0.7 to 1.7±1.5; 17.1%–63.0%).10 29

Even though we determined that drugs from the central nervous system are the most common drugs identified with the German ACB score and the second most common for ADS, our top 10 anticholinergic drugs showed a more diverse spectrum of drugs. Particularly, drugs with low to moderate anticholinergic effects occurred in our top 10 list for both scores. As 80.2% of the anticholinergic drugs according to ADS and 73.4% of the anticholinergic drugs according to German ACB are anticholinergics with a score of 1, it is important to also focus on the drugs with only mild anticholinergic potential during prescribing and reviewing patients’ medications. Furthermore, drugs treating cardiovascular conditions highly contributed to the level 1 anticholinergic drugs in both scores. However, in multimorbid elderly patients, it is common to coprescribe drugs like furosemide, triamterene, digoxin and captopril to treat multiple conditions.32 33 It is stated that especially the cumulative anticholinergic effect contributes to higher anticholinergic scores and even leads to hospitalisation and higher risk for mortality.34–36 A lot of the mentioned and detected level 1 drugs are peripherally acting anticholinergic drugs. However, as the permeability of the blood–brain barrier is increased and at the same time the P-glycoprotein function is decreased with growing age, elderly people are more vulnerable towards anticholinergic ADRs.5

---

**Table 3** The influence of age, sex and the number of taken drugs on the anticholinergic drug use according to anticholinergic drug scale (ADS) score in multimorbid elderly patients (significant p values are marked in bold)

<table>
<thead>
<tr>
<th>ADS score</th>
<th>Number of patients</th>
<th>Mean</th>
<th>SD</th>
<th>Range</th>
<th>95% CI</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;80 years</td>
<td>2635</td>
<td>0.71</td>
<td>1.24</td>
<td>0–11</td>
<td>0.67 to 0.76</td>
<td></td>
</tr>
<tr>
<td>≥80 years</td>
<td>554</td>
<td>0.91</td>
<td>1.38</td>
<td>0–9</td>
<td>0.79 to 1.02</td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>1298</td>
<td>0.65</td>
<td>1.15</td>
<td>0–8</td>
<td>0.58 to 0.71</td>
<td>p=0.001</td>
</tr>
<tr>
<td>Female</td>
<td>1891</td>
<td>0.82</td>
<td>1.34</td>
<td>0–11</td>
<td>0.76 to 0.86</td>
<td></td>
</tr>
<tr>
<td>0–7 drugs</td>
<td>1688</td>
<td>0.36</td>
<td>0.82</td>
<td>0–9</td>
<td>0.33 to 0.4</td>
<td>p&lt;0.001</td>
</tr>
<tr>
<td>8–29 drugs</td>
<td>1501</td>
<td>1.18</td>
<td>1.52</td>
<td>0–11</td>
<td>1.1 to 1.25</td>
<td>p&lt;0.001</td>
</tr>
</tbody>
</table>

**Table 4** The influence of age, sex and the number of taken drugs on the anticholinergic drug use according to German anticholinergic burden (ACB) score in multimorbid elderly patients (significant p values are marked in bold)

<table>
<thead>
<tr>
<th>ACB score</th>
<th>Number of patients</th>
<th>Mean</th>
<th>SD</th>
<th>Range</th>
<th>95% CI</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;80 years</td>
<td>2635</td>
<td>1.18</td>
<td>1.61</td>
<td>0–11</td>
<td>1.12 to 1.24</td>
<td></td>
</tr>
<tr>
<td>≥80 years</td>
<td>554</td>
<td>1.27</td>
<td>1.62</td>
<td>0–9</td>
<td>1.14 to 1.41</td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>1298</td>
<td>1.04</td>
<td>1.42</td>
<td>0–9</td>
<td>0.96 to 1.11</td>
<td>p=0.1992</td>
</tr>
<tr>
<td>Female</td>
<td>1891</td>
<td>1.30</td>
<td>1.73</td>
<td>0–11</td>
<td>1.22 to 1.38</td>
<td></td>
</tr>
<tr>
<td>0–7 drugs</td>
<td>1688</td>
<td>0.60</td>
<td>1.02</td>
<td>0–9</td>
<td>0.56 to 0.65</td>
<td>p&lt;0.001</td>
</tr>
<tr>
<td>8–29 drugs</td>
<td>1501</td>
<td>1.86</td>
<td>1.87</td>
<td>0–11</td>
<td>1.76 to 1.95</td>
<td>p&lt;0.001</td>
</tr>
</tbody>
</table>
Particularly multimorbid elderly patients are vulnerable for polypharmacy. So it is not surprising that we identified polypharmacy as risk factor for a high German ACB and ADS score. Besides this, we detected that female patients seem to be more vulnerable towards the exposure with anticholinergic drugs. This gender shift was also observed in several studies, although most studies used different tools to identify the anticholinergic burden. The increased vulnerability of women towards anticholinergic drug exposure might be explainable by the fact that women have a higher health awareness than men. In addition, rates of depression are higher in the female population, and we identified drugs from the central nervous system as one of our largest drug groups contributing to the anticholinergic burden.

**Association of anticholinergic drug use with the cognitive function**

Multivariate analysis revealed that a higher anticholinergic burden according to ADS and also the German ACB score is associated with a decreased cognitive function according to an increasing poorly performance in the LDST. Interestingly, the newly developed German ACB score showed similar results in our adjusted model in comparison with the already well-established ADS score. A lot of studies prove that a high anticholinergic burden is associated with a decreased cognitive function as well. However, there are also opposite findings. For example, Kersten et al could determine that there was no association between cognitive impairments and anticholinergic drug use according to ADS score.

The differences in the outcomes might be explained by several factors. First, it is sometimes difficult to show a homogenous association between anticholinergic drug use and cognitive function, because there is a broad heterogeneity in cholinergic brain reserve in each individual that leads to differences in the sensitivity to central anticholinergic effects, and second, the used tools for detecting anticholinergic drugs and drugs with anticholinergic activity and measuring the cognitive function of the patients differs between the studies and not always fits the country-specific prescribing habits. However, Gray et al detected in a prospective cohort study over a time period of 7.3 years that 23% of the patients 65 years old and older develop a dementia and thereof 80% used anticholinergic drugs. As already mentioned, patients were excluded from MultiCare study when they were diagnosed with dementia and/or were living in nursing homes. Even though there was no standardised tool for diagnosing dementia due to the different GPs in the different study centres, we can assume that our patient collective had less cognitive impairments than the collectives in other studies. So it is quite interesting that our patient collective already shows an association between decreased cognitive function based on poorer results in LDST and a high anticholinergic score. That demonstrates the importance of rational prescribing and also deprescribing, even in presumed healthier elderly patients. Drugs with anticholinergic activity are widely prescribed, but we need to evaluate the pros and cons of their usage. On the
One hand, alternative treatments are partly not available or appropriate, and on the other hand, there are risk of anticholinergic side effects. That is why we need to weigh the risk between deprescribing and a possible undertreatment of critical conditions. Consequently, we are in need to develop interdisciplinary processes for deprescribing. Ailabouni et al invented a five-step systematic intervention in deprescribing anticholinergic and sedative drugs for a small patient collective. Although they could not report an improvement in cognitive function over a time period of 6 months after deprescribing, they could lower the used medication in mean about 2.1 drugs per patient. They also detected that patients reported significantly less adverse effects, reduced falls and mortality. However, for deprescribing, we are in need for validated tools, with regard to anticholinergic drug use and regarding potentially inappropriate medication. In addition, it is interesting to know whether it is necessary to evaluate patients’ medication concerning PIM lists and anticholinergic burden lists. Studies revealed a high proportion of anticholinergics and sedatives within the detected PIM, but there was no analysis with regard to the necessity of using PIM tools and tools to evaluate the anticholinergic burden.

Although we determined by including FORTA PIM into the regression model a decrease of the regression coefficient for ADS and German ACB score about −0.1, the anticholinergic scores and therefore the use of anticholinergic drugs according to ADS and German ACB score still seemed to have a negative influence on the cognitive function on multimorbid elderly patients. In addition, the FORTA list did not cover all drugs detected with ADS and/or German ACB score. So, we assume that multimorbid elderly patients could benefit from the use of both lists (PIM and anticholinergic burden).

**Strength and limitations**

Our study has some strengths and limitations. Unfortunately, we could not underline our results by showing an impact of anticholinergic drug use on peripheral anticholinergic ADR (eg, dry mouth, blurred vision, constipation, nausea, urinary retention, impaired sweating or...
tachycardia) because such parameters were not gained during data collection. However, other studies showed that anticholinergic drug use is associated with significantly increased mouth dryness.31

Most studies had a less healthy patient collective than we had, due to the fact that we excluded patients living in nursing homes. De Vreese et al detected that especially patients from nursing homes are at a greater risk of receiving anticholinergic drugs. However, we could demonstrate that even the apparently more healthy elderly patients are in great risk for receiving anticholinergic drugs and thereby suffering from anticholinergic side effects in association with decreased cognitive function. Moreover, we could not evaluate the length of intake of anticholinergic drugs. Further studies, especially longitudinal studies, are necessary to evaluate the decrease in cognitive function over time. As cognitive impairments is a complex clinical symptom and the LDST only addresses one single aspect of cognition, further tests would help to underline our findings. However, a strength of our study is that we performed a multivariate analysis, including among other number of diseases weighted by severity. A sensitivity analysis was performed, revealing that FORTA PIM has to be included as a confounder in the regression model. In contrast to the number of used drugs, which had no significant influence on the results in LDST (ADS score: p=0.257, regression coefficient: –0.04; ACB score: p=0.518; regression coefficient: −0.02). An additional strength is also the advanced treating of missing values via hot deck imputation.

Taken together, anticholinergic drugs and drugs with anticholinergic activity in multimorbid elderly adults appear to be associated with harms that, in certain circumstances, outweigh their potential benefit. We could determine that a high anticholinergic score is associated with a reduced cognitive function, according to increased poorer results in LDST, in multimorbid elderly patients. In addition, we showed that especially drugs with low anticholinergic risk, for example, for treating cardiovascular conditions, contribute to the anticholinergic burden.

Further studies are needed, especially showing the effect on patient outcome on deprescribing anticholinergic drugs over a longer time period and longitudinal studies to demonstrate the development of cognitive function under use of anticholinergic drugs over time.

In summary, a high anticholinergic burden and therefore anticholinergic drug use is associated with a decreased cognitive function in multimorbid elderly patients. In order to contribute to an improvement in drug therapy safety, we need to invent strategies for rational prescribing and deprescribing.

CONCLUSION

Our study demonstrated that it is important to gain greater awareness for the risk of using anticholinergic drugs in multimorbid elderly patients and that there exist tools that are easy to use in medical routine to calculate the anticholinergic burden of this vulnerable patient group. Furthermore, we pointed out that the newly invented German ACB score by Kiesel et al seems to generate comparable results with already validated and established tools. However, it needs to be validated in future in order to gain data about the safe use of this tool.

As shown in our study, it is also important to question lower potential anticholinergic drugs, since cumulative effects of those low potential anticholinergic drugs can lead to high anticholinergic burdens as well.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available on reasonable request. The data that support the findings of this study are available from Professor Hendrik van den Bussche, but restrictions apply to the availability of these data, which were used under licence for the current study, and so are not publicly available. Data are however available from the authors on reasonable request and with permission of Professor Hendrik van den Bussche.

Supplemental material This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been peer-reviewed. Any opinions or recommendations discussed are solely those of the author(s) and are not endorsed by BMJ. BMJ disclaims all liability and responsibility arising from any reliance placed on the content. Where the content includes any translated material, BMJ does not warrant the accuracy and reliability of the translations (including but not limited to local regulations, clinical guidelines, terminology, drug names and drug dosages), and is not responsible for any error and/or omissions arising from translation and adaptation or otherwise.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

ORCID iDs
Caroline Krüger http://orcid.org/0000-0001-9080-1562
Ingrid Schäfer http://orcid.org/0000-0002-1038-7478

REFERENCES
20 Ms S, S. Tsai, H, Tsai, P. N. Comparison of anticholinergic risk scales and associations with adverse health outcomes in older people [Internet]. J Am Geriatr Soc 2015.
22 Kiesel EK, Hopf YM, Drey M. An anticholinergic burden score for German prescribers: score development. BMC Geriatr 2018;18:239.