Non-persistence and non-adherence to MTX therapy in patients with rheumatoid arthritis: a retrospective cohort study based on German RA patients

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Objective: This study aimed to assess the level of nonpersistence (NP) and nonadherence (NA) to methotrexate (MTX) therapy in German patients with rheumatoid arthritis (RA).

Materials and methods: Based on German claims data, RA patients who received a MTX therapy (subgroup: treatment-naive patients) were analyzed. NP was defined as treatment gap >12 weeks. Regarding NA, it is the overall medication possession ratio (MPR) during an observational period of 12 or 24 months after therapy, and the MPR is calculated only for the periods of therapy continuation; NA was defined as MPR <80%.

Results: A total of 7,146 RA patients who received at least one MTX prescription (subgroup: 1,211 treatment-naive patients) could be observed (mean age: 64.4 years, 73.6% female). Percentage of NP patients among MTX-naive patients after 6, 12 and 18 months was 16.7%, 34.0% and 36.7%, respectively. After MTX therapy discontinuation, 39.9% had restarted their MTX therapy, 13.8% had received another non-MTX synthetic disease-modifying antirheumatic drug (sDMARD), 8.1% had biological DMARD (bDMARD) and 49.2% had not received any DMARD prescription at all. Overall, 12- and 24-month MPRs for MTX therapy were 83.0% and 76.5% with a percentage of NA patients of 25.8% and 33.8%, respectively.

Conclusion: NP to MTX treatment seems to be common in one-fourth of German patients with RA. An additional number of patients, at least 6.5%, are also affected by NA. A considerable percentage of RA patients who discontinued MTX therapy do not receive any follow-up DMARD therapy.

Keywords: rheumatoid arthritis, RA, MTX therapy, adherence to MTX therapy, persistence with MTX therapy, discontinuation of MTX therapy

Introduction

Rheumatoid arthritis (RA) is a chronic systemic autoimmune inflammatory disease with a prevalence of up to 1% in western countries.1,2 Poorly controlled RA results in severe progressive joint damage, functional disability, morbidity, poor health-related quality of life and higher mortality.3,4

According to current treatment guidelines, methotrexate (MTX) as a synthetic disease-modifying antirheumatic drug (sDMARD) is the appropriate first-line agent (in combination with glucocorticoids) for the majority of RA patients as part of a treat-to-target approach.5 If MTX therapy fails to improve symptoms after 3 months or fails to reach treatment targets after 6 months, it is recommended to move to another
sDMARD, another sDMARD combination or to add a biological DMARD (bDMARD).

When taken as prescribed, sDMARD and bDMARD therapy can result in remission of RA disease activity.\textsuperscript{3} However, nonadherence (NA), defined as poor implementation of a generally continued therapy, and nonpersistence (NP), defined as general discontinuation of therapy, have been reported to be major challenges in the real-world treatment of RA patients.\textsuperscript{6} Although bDMARD therapy is currently used in the treatment of a minority of RA patients,\textsuperscript{7} more is known about persistence with/adherence to bDMARD therapy than persistence with/adherence to first-line sDMARD and, specifically, MTX therapy.\textsuperscript{8–10} Furthermore, existing studies addressing NA or NP associated with MTX therapy in RA patients failed to differentiate between persistence and adherence.\textsuperscript{6,11}

Consequently, the main objectives of this study were to assess the level of NP and NA in German patients with RA and to describe the DMARD treatment of MTX discontinuers.

Materials and methods

Data source and patient sample

We did a retrospective noninterventional cohort analysis based on anonymized claims data provided by a German statutory health care fund (AOK PLUS; 2.8 million insured) for the years 2010–2013. The analysis included all RA patients permanently enrolled in the database from January 01, 2010, until the end of the observational period; death during the observational period was the only exception to the disrupted enrollment requirement. A patient was defined as RA patient if at least one outpatient RA-ICD-10 diagnosis (ICD-10 M05: seropositive RA; M06: other RA; excluding M06.1: adult-onset still disease) and/or at least one inpatient RA-ICD-10 diagnosis was documented prior to the first observed reference prescription of study medication.

Observational periods

We analyzed persistence with/adherence to MTX therapy in common use (Anatomical-Therapeutic-Chemical [ATC] codes M01CX01/L04AX03). Analyses were carried out in all patients receiving MTX therapy, defined as at least one observed MTX prescription from January 01, 2010 to December 31, 2011, with a potential follow-up period of 24 months. Analyses also included a subgroup of MTX treatment-naive patients, defined as no MTX prescription in the 12 months prior to the first observed MTX prescription in the period between January 01, 2011 and December 31, 2011. Reference period, inclusion period, index date and follow-up period are explained in Figure 1. In all the analyses, patients who received any MTX therapy as intravenous infusion therapy were excluded, because no reliable assumptions could be made regarding the prescribed daily dosage (PDD).

Assessment of treatment persistence

Persistence with MTX treatment was reported for the following groups: all treatment-naive patients and, as subgroups, MTX treatment-naive patients who received 1) MTX monotherapy only (no concomitant treatment with other DMARD agents until the end of the follow-up period or the first observed treatment gap, whichever came first), 2) a combination therapy of MTX and other sDMARDs (sDMARD agents: leflunomide/ciclosporin/azathioprine/sodium aurothiomalate/sulfasalazin/metronidazole/tinidazole; ATC codes: L04AA13/L04AD01/L04AX01/M01CB01/M01CX02/P01BA01/P01BA02), 3) a combination therapy of MTX and bDMARDs (bDMARD agents: anakinra/tocilizumab/abatacept/adalimumab/certolizumab/etanercept/golimumab/infliximab; ATC codes: L04AC03/L04AC07/L04AA24/L04AB04/L04AB05/L04AB01/L04AB06/L04AB02), 4) oral MTX therapy only (with and without concomitant DMARD/bDMARD medication), 5) subcutaneous MTX therapy only (with and without concomitant DMARD/bDMARD medication) and finally 6) a change of application mode from subcutaneous to oral MTX therapy or vice versa (with and without concomitant DMARD/bDMARD medication) during the follow-up period. We allowed for any previous non-MTX medication and any concomitant medication during MTX therapy.

Our analysis was based on the number of days covered by observed prescriptions recorded in the database. Because data on PDD were missing, which is common in most known claims datasets, we developed and validated a model providing assumptions for PDD. This model was developed for 12 different strengths and three different pack sizes per strength for MTX therapy as available in the German pharmaceutical market.

We assumed the following: the PDD associated with an MTX prescription was assumed to be equal to one tablet/subcutaneous injection a week. Stockpiling was included by assuming that, in case there were overlapping medications, the previous supply was taken fully before the new supply was initiated. Additionally, hospitalization periods were excluded from observed days because drug supply during hospitalization was assumed to be arranged by hospitals.

However, in case a change in the application mode was observed (from oral to subcutaneous or vice versa), we assumed that stock of the previous prescription would no longer be used. Furthermore, we applied two additional
assumptions. First, in case the dosage strength of a second prescribed MTX package was different from the previous one (ie, 2.5 mg instead of 10 mg), we observed the third follow-up prescription. If that included the dosage of the first observed prescription again (ie, 10 mg) or a dosage resulting from the sum of both previous prescribed dosages (ie, 12.5 mg), we assumed the second one to be an add-on dosage to the first prescribed dosage, leading to a PDD, which was the sum of both first prescriptions (ie, 12.5 mg/wk). However, if the third prescription included the same strength as the second prescription (ie, third prescription with 2.5 mg/wk again), we assumed that there was a change in prescribed PDD (ie, 2.5 mg/wk).

Second, we calculated on a continuous basis the resulting medication possession ratios (MPRs) for three subsequent MTX prescriptions. If the resulting MPR was &gt;200%, we assumed the PDD to be twice as high as the originally assumed PDD for the period covered by these three prescriptions for the respective period.

It should be noted that the PDD was reassessed with each observed MTX prescription. Both abovementioned specific assumptions consequently might have led to revised PDDs, but this only applied to a period from the currently observed prescription until the next one. With the day of the follow-up prescription, a new assessment of the PDD including both assumptions was done.

The developed PDD model as outlined previously was subsequently validated in a separate validation study. Based on a medical chart review, five study sites documented 934 MTX prescriptions and the associated, physician-based PDDs of 160 RA patients (inclusion criteria: age above 18 years, confirmed RA diagnosis and current/previous treatment with MTX for at least 24 months), using predefined case report forms. Specifically, we asked study sites to document all MTX prescriptions (including prescription date, application mode, dosage strength, package size and recommended dosage) of the last 2 years for each patient. Based on these documentations, we calculated the overall MPR between the first and the last documented MTX prescriptions, as defined subsequently (ratio of the supplied days and the duration between the first and the last documented prescriptions). The percentage of patients with an MPR &lt;80% based on rheumatologists’ data was compared with the percentage of patients with MPR &lt;80% based on different PDD models.
as used in the main claims-based data analysis (Table 1). The PDD model with the lowest deviation between rheumatologists’ records and our PDD assumptions was used as base case in the full claims data analysis; it was the model described previously.

NP was defined as a treatment gap of >12 weeks in the base case; in a sensitivity analysis, we also used 24 weeks as threshold. Persistence was reported based on Kaplan–Meier (KM) curves and the percentage of patients who could be classified as NP at 6, 12 and 18 months after therapy start (index date). Furthermore, we reported the average treatment continuation time for the entire period of 18 months and time until MTX treatment discontinuation for patients who discontinued MTX therapy after 6, 12 and 18 months.

**Assessment of postdiscontinuation treatment**

In our base case scenario (12-week gap, NP assessment 18 months after index date), we reported whether patients who discontinued MTX treatment received any follow-up DMARD treatment during 12 months after treatment discontinuation and, if applicable, which treatment they received. Patients who could not be observed for a 12-month follow-up after an NP event were excluded from this analysis.

**Assessment of treatment adherence**

Treatment adherence was analyzed in two ways. First, for the overall sample, including those patients who may have discontinued therapy during the follow-up period and those continuing their therapy, we analyzed the overall MPR for both 1 and 2 years based on the following formula:

\[
\text{MPR} = \frac{\text{Number of days supply received during observational period}}{(365 \text{ OR } 730 \text{ days} - \text{hospitalization days} - \text{days after death})}
\]

**Table 1** Applied PDD models: results of the validation study

<table>
<thead>
<tr>
<th>Model assumptions</th>
<th>Base case scenario</th>
<th>Scenario 1</th>
<th>Scenario 2</th>
<th>Scenario 3</th>
<th>Scenario 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDD assumed to be equal to one pill/injection per week</td>
<td>Included</td>
<td>Included</td>
<td>Included</td>
<td>Included</td>
<td>Not included</td>
</tr>
<tr>
<td>Hospitalization periods taken out from observation</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Stockpiling if no change of application mode was observed</td>
<td>Included</td>
<td>Included</td>
<td>Included</td>
<td>Included</td>
<td>Included</td>
</tr>
<tr>
<td>Consideration of possible dosage strength combination (eg, one pill of 10 mg and one pill of 2.5 mg for a prescribed dosage of 12.5 mg/wk)</td>
<td>Included</td>
<td>Not included</td>
<td>Not included</td>
<td>Not included</td>
<td>Not included</td>
</tr>
<tr>
<td>PDD assumed to be 200% of the originally assumed PDD if the continuously calculated MPR for three subsequent prescriptions was &gt;200%</td>
<td>Included</td>
<td>Not included</td>
<td>Included</td>
<td>Not included</td>
<td>Not included</td>
</tr>
<tr>
<td>Mean MPR for 160 RA validation study patients based on above model</td>
<td>85.78</td>
<td>85.82</td>
<td>86.30</td>
<td>86.35</td>
<td>66.09</td>
</tr>
<tr>
<td>Mean MPR for 160 RA validation study patients based on reported PDD by rheumatologists</td>
<td>82.11 %</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deviation in MPR between PDD model and reported PDD by rheumatologists (percentage points)</td>
<td>+3.66</td>
<td>+3.71</td>
<td>+4.19</td>
<td>+4.24</td>
<td>-16.02</td>
</tr>
<tr>
<td>Percentage of RA validation study patients with MPR &lt;80% based on above model</td>
<td>26.88</td>
<td>26.88</td>
<td>26.25</td>
<td>26.25</td>
<td>73.13</td>
</tr>
<tr>
<td>Percentage of RA validation study patients with MPR &lt;80% based on PDD reported by rheumatologists</td>
<td>35.63</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deviation in percentage of RA patients between PDD model and reported PDD by rheumatologists (percentage points)</td>
<td>-8.75</td>
<td>-8.75</td>
<td>-9.38</td>
<td>-9.38</td>
<td>+37.50</td>
</tr>
</tbody>
</table>

**Note:** PDD assumed to be equal to the DDD (as defined by WHO) of the last prescription.

**Abbreviations:** PDD, prescribed daily dosage; DDD, defined daily dosage; MPR, medication possession ratio; RA, rheumatoid arthritis.
In our second NA analysis, we explored adherence only for the period in which a patient continued MTX therapy (no treatment gap > 12 weeks). If a treatment gap of > 12 weeks was observed, the observational period for this specific patient lasted only from index date until the day of the last prescription before the first treatment gap > 12 weeks was observed (Figure 1):

\[ \text{MPR} = \frac{\text{Number of days supply received during persistent period}}{\text{(Days between first and last prescription without any treatment gaps > 12 weeks – hospitalization days)}} \]

Adherence was reported for all RA patients who received MTX treatment and, as subgroups, MTX treatment-naive patients, patients who received MTX monotherapy, oral MTX therapy or subcutaneous MTX therapy or who experienced a switch in application form.

Adherence was reported in the following three ways:
1. as mean/median MPR,
2. as the percentage of patients with MPR < 80%,
3. in a sensitivity analysis, as the percentage of patients with MPR < 70% < 90%.

In case an MPR was above 100%, it was truncated to 100%.

Statistical analysis
Significance of differences between discontinuation rates as shown by KM curves was tested by means of log-rank tests. Significant level was set at \( P < 0.05 \), and all reported \( P \)-values were two sided. Descriptive evaluations were executed with Microsoft SQL Server 2008 and Microsoft Excel 2010. All other statistical analyses were performed with STATA 13.1.

Ethics and scientific approval
This was a noninterventional, retrospective study analyzing anonymized data as defined by section 3 (6) of the Federal Data Protection Act.\textsuperscript{12} For the first part of the study, which was a medical chart review and for the subsequent secondary data analysis of claims data, anonymized data were only used. Thus, ethical approval and informed consent from patients were not obtained. This is in accordance with national guidelines and recommendations,\textsuperscript{13,14} the Federal Data Protection Act (the use of anonymized data is not restricted by means of data protection regulations)\textsuperscript{12} and with the policy of the institutions conducting the analysis (IPAM and AOK PLUS).

However, the study was evaluated by a scientific steering committee to which all the authors belonged.

Results
MTX-experienced and MTX-naive RA samples
In our sample, 7,146 RA patients who received at least one MTX prescription in 2010 or 2011 (mean age: 64.4 years, 73.6% female) were observed. Of these, 1,211 started MTX treatment in 2011 (mean age: 62.2 years, 71.5% female).

Patient characteristics are listed in Table 2.

For MTX-experienced and MTX-naive patients, percentages of patients who received MTX monotherapy (no concomitant sDMARD/bDMARD therapy) were 79.3% and 83.7%, respectively. Percentages of RA patients who received an MTX+sDMARD combination therapy (without any bDMARD) were 10.4% and 9.7%, and percentages of RA patients who received at least one bDMARD prescription during follow-up were 10.3% and 6.6%, respectively. In all, 71.5% and 71.1% of MTX-experienced and MTX-treatment-naive patients received oral MTX therapy only, 18.5% and 16.9% received subcutaneous MTX therapy only and 10.0% and 12.0% switched between application modes during the follow-up period.

Assessment of treatment persistence
The results of the MTX treatment persistence analysis based on 1,211 RA patients who started MTX treatment are presented in Figure 2A and B and Table 3. Based on a > 12-week treatment gap definition, percentages of NP patients after 6, 12 and 18 months were 16.7%, 34.0% and 36.7%, respectively. Mean time until discontinuation was 11.9, 22.2 and 25.2 weeks, respectively. The estimated mean overall time on treatment within the 18-month follow-up period was 58.4 weeks. Based on all patients showing NP after 6 months, 42.4% received a single MTX-index prescription (early NP).

If patients receiving MTX monotherapy were compared with patients receiving a combination therapy of MTX and other sDMARDs/bDMARDs, persistence was not significantly different (Figure 2A). Additionally, persistence rates did not differ between different types of application modes (Figure 2B). A change in application was associated with a higher treatment continuation rate. However, since a switch itself is associated with at least two different MTX prescriptions, this obviously excluded all NP patients with only one prescription from this group.
Assessment of postdiscontinuation treatment

Based on the base case scenario (12-week gap, NP assessment 18 months after index date), 36.7% of the observed MTX-naive patients had discontinued their treatment. After discontinuation, 419 patients (94.4% of all NP patients) could be observed for a follow-up period of 12 months after the NP event. Of these, 39.9% restarted their MTX therapy (patients with/without at least one rheumatologist visit in the follow-up period: 43.8/36.2%; \( P=0.070 \)), 13.8% received another non-MTX sDMARD therapy (with/without rheumatologist visit: 15.4/12.4%; \( P=0.368 \)), 8.1% received at least one bDMARD prescription (with/without rheumatologist visit: 12.4/4.1%; \( P=0.002 \)) and 49.2% did not receive any DMARD prescription at all (with/without rheumatologist visit: 43.3/54.6%; \( P=0.021 \)).

Assessment of treatment adherence

The overall 12- and 24-month MPRs for MTX therapy in the analyzed MTX treatment-experienced RA patients were 83.0% and 76.5%, respectively. The percentage of NA patients (MPR <80%) was 25.8% for the 12-month follow-up and 33.8% for the 24-month follow-up (Table 4).

In our second adherence analysis, which was carried out during periods of general treatment continuation only (no gaps >12 weeks), mean MPRs were 94.7% (all patients), 95.0% (treatment-naive patients only), 94.8% (MTX monotherapy), 94.5% (MTX plus other DMARD therapy), 93.7% (MTX+bDMARD therapy), 96.0% (oral MTX therapy), 91.6% (subcutaneous MTX therapy) and 92.9% (change between oral and subcutaneous therapy).
Table 4 shows the corresponding percentages of patients with MPR <80% and the results of scenario analyses (MPR thresholds 70% or 90%). The distribution of MPRs across our patient sample is shown in Figure S1.

**Discussion**

**Study objectives and main results**

Based on a German claims dataset, we aimed to assess the level of persistence and adherence to MTX therapy in RA.
Table 3 Results of medication – persistence analysis (n=1,211 RA patients who started MTX therapy)

<table>
<thead>
<tr>
<th>Treatment persistence</th>
<th>After 6 months</th>
<th>After 12 months</th>
<th>After 18 months</th>
<th>After 6 months</th>
<th>After 12 months</th>
<th>After 18 months</th>
</tr>
</thead>
<tbody>
<tr>
<td>NP definition: gap of at least 12 weeks/24 weeks</td>
<td>12 weeks</td>
<td>12 weeks</td>
<td>12 weeks</td>
<td>24 weeks</td>
<td>24 weeks</td>
<td>24 weeks</td>
</tr>
<tr>
<td>Percentage of all MTX treatment-naive patients (n=1,211) who discontinued MTX therapy</td>
<td>16.7</td>
<td>34.0</td>
<td>36.7</td>
<td>11.2</td>
<td>25.1</td>
<td>26.5</td>
</tr>
<tr>
<td>Mean time until discontinuation of MTX therapy for abovementioned patients (median)</td>
<td>11.9 weeks (10 weeks)</td>
<td>22.2 weeks (27 weeks)</td>
<td>25.2 weeks (30 weeks)</td>
<td>11.2 weeks (10 weeks)</td>
<td>22.7 weeks (30 weeks)</td>
<td>24.8 weeks (30 weeks)</td>
</tr>
<tr>
<td>Estimated mean overall treatment continuation time$^c$</td>
<td>58.4 weeks</td>
<td>63.7 weeks</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 4 Results of medication – adherence analysis (all RA patients who received MTX therapy)

<table>
<thead>
<tr>
<th>Base case: from first until last observed MTX prescription or until first day of an observed treatment gap (&gt;12 weeks); whichever came first</th>
<th>Scenario 1: interval-based calculation: 12 months since index prescription</th>
<th>Scenario 2: interval-based calculation: 24 months since index prescription</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>6,104$^a$</td>
<td>7,146$^b$</td>
</tr>
<tr>
<td>Mean MPR (%) (median)</td>
<td>94.7 (97.7)</td>
<td>83.0 (100.0)</td>
</tr>
<tr>
<td>Percentage of patients with an MPR &lt;80%</td>
<td>6.5</td>
<td>25.8</td>
</tr>
<tr>
<td>Percentage of patients with an MPR &lt;70%</td>
<td>2.5</td>
<td>22.6</td>
</tr>
<tr>
<td>Percentage of patients with an MPR &lt;90%</td>
<td>18.3</td>
<td>29.1</td>
</tr>
<tr>
<td>Percentage of following patient subgroups with an MPR &lt;80%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• MTX treatment-naive patients</td>
<td>6.3</td>
<td>41.8</td>
</tr>
<tr>
<td>• MTX mono</td>
<td>6.8</td>
<td>23.6</td>
</tr>
<tr>
<td>• MTX+bDMARD</td>
<td>6.8</td>
<td>14.9</td>
</tr>
<tr>
<td>• MTX+other DMARD</td>
<td>8.7</td>
<td>12.8</td>
</tr>
<tr>
<td>• Oral MTX</td>
<td>5.1</td>
<td>25.2</td>
</tr>
<tr>
<td>• Subcutaneous MTX</td>
<td>12.1</td>
<td>30.2</td>
</tr>
<tr>
<td>• Change in MTX application mode</td>
<td>8.4</td>
<td>19.2</td>
</tr>
</tbody>
</table>

Notes: $^a$Estimated survival time based on KM estimates. Note that the KM estimates did not represent the median survival time.

Abbreviations: NP, nonpersistence; MTX, methotrexate; DMARD, disease-modifying antirheumatic drug; bDMARD, biological DMARD; RA, rheumatoid arthritis; KM, Kaplan–Meier.

patients and to identify the treatment patterns of RA patients after discontinuation of MTX therapy.

The strength of this study is its use of a very large observational database. Additionally, we differentiated between treatment persistence and treatment adherence since we believe that these are different real-world phenomena that may be caused by different factors and may lead to different conclusions. For example, discontinuation of MTX therapy (NP) may be caused by patients’ behavior, but it may also be based on a physician’s decision not to prescribe MTX anymore because of severe safety concerns and/or treatment failure. NA, on the other hand, happens despite existing prescriptions of a treating physician and can be assumed to be caused mainly by patient-centered factors.$^{15–17}$ However,
in most publications evaluated by the authors of this study, adherence and persistence are analyzed jointly, so that reported percentages of NA patients still include patients discontinuing their therapy during a predefined observational period. If, for example, and based on our database, the percentage of NA patients is calculated as MPR <80% for a follow-up of 2 years irrespective of continuation/ discontinuation of therapy during that period, then 33.8% of the patients need to be classified as NA. However, most of these NA patients discontinued treatment instead of implementing their therapy in a suboptimal way. So, in our main analysis, the percentage of NA patients decreased to 6.5%.

The only obvious disadvantage of our methodology in comparison to the more general MPR approach is that we may have underestimated NA for NP patients, because periods after a restart of MTX therapy, which is also potentially vulnerable to NA, were not included in our adherence analysis.

Our study, using a gap >12 weeks’ threshold, revealed that 34.0% of MTX treatment-naive patients discontinued their MTX therapy after 12 months. Considering an additional 6 months and observing 18 months after therapy initiation, we found that only an additional 2.7% of MTX patients discontinued therapy. This may indicate that persistence is a critical issue at the beginning of that therapy. It is known from clinical studies that this may be explained by severe side effects/toxicity, by low therapy effectiveness in specific RA patients or by patient’s decisions. The percentage of RA patients who poorly implemented MTX therapy, defined as NA associated with an MPR <80% during periods of general MTX therapy discontinuation, was 6.5%, indicating a rather good quality of therapy implementation during its general continuation.

So, early MTX therapy discontinuation seems to be the most critical issue, based on our observations. Whether an observed early noneffectiveness of therapy, severe side effects, a poor management of light side effects or even other reasons explain this therapy discontinuation was not explored in our study. If there would be potential for a more closer and better management of RA patients in their early phase of MTX treatment, this could both improve MTX therapy outcomes and could reduce the percentage of patients in need of therapy intensification, eg, with bDMARDs.

A systematic international review summarizing 35 articles (1985–2008) that addressed adherence to/persistence with DMARD treatment in RA patients (of these, 10 articles reported adherence rates) reported a percentage of 30–100 of RA patients to be adherent to DMARD therapy. The substantial difference in reported NA rates in these publications is mainly due to different methodologies used in the various analyses to measure adherence: these range from electronic event monitoring systems to pill counts, self-reports and claims data analyses.

Previous US claims data analyses reported numbers of NP patients similar to our analysis, if methodological differences are taken into account. In line with that, a recent analysis of a Danish nationwide longitudinal cohort of patients hospitalized for RA reported a substantial percentage of >10% of patients to be early MTX therapy discontinuers, defined as no second filled MTX prescription.

However, we report a lower percentage of patients being nonadherent to MTX therapy. This is mainly because we analyzed adherence during periods of MTX treatment continuation only. If both persistence and adherence are analyzed jointly, nearly 40% of RA patients face either NP or NA, which is in accordance with previous publications, also in a comparison to other long-term chronic diseases.

Due to the nature of our dataset, we were able to describe the drug treatment of RA patients after MTX therapy discontinuation. We showed that 39.9% of patients restarted MTX therapy after a gap >12 weeks and that follow-up prescriptions of another DMARD (13.8%) or adding a bDMARD (as monotherapy or combination therapy; 8.1%) were common. Nevertheless, a substantial percentage of patients (49.2%) did not receive any DMARD treatment in the 12 months after MTX therapy discontinuation, which is not in line with treatment guidelines. One reason could be that our patients started MTX therapy but were misdiagnosed with RA. Due to the nature of our dataset, we could not assess in which percentage of NP patients this was the case.

Additionally, we found that the percentage of NP patients who did not receive any DMARD follow-up treatment was substantially higher in patients who did not visit a rheumatologist in the year after MTX discontinuation. This was the case in 52.0% of our observed patients who discontinued MTX therapy.

Limitations
We acknowledge some limitations of our analysis. Due to longitudinal limitations of our dataset at the time of analyses, we could only observe 24 months after the start of MTX therapy. Further research is needed to describe longer term persistence/adherence in RA patients who started MTX therapy. In addition to that, generally, we did not explore any causes of NP or NA to MTX therapy in our study because of lack of data.
We defined NP as a treatment gap >12 weeks and NA as an MPR <80%. While these thresholds are widely used in general adherence/persistence literature and in claims data analyses addressing RA patients, they have so far hardly been clinically validated.22,29 We dealt with this weakness by reporting results of a sensitivity analysis using a gap >24 weeks as NP definition and by using different MPR thresholds of 70%, 80% and 90%.

Finally, we only analyzed filled prescriptions in our database. We may have underestimated NA and NP because specific patients may have filled prescriptions but did not take them as prescribed.

Conclusion and implications for practice
NP to MTX treatment appears to be common in one-fourth of German patients with RA. An additional percentage of at least 6.5% of patients is also affected by NA. A considerable percentage of RA patients discontinuing MTX therapy do not receive any follow-up DMARD therapy. Consequently, efforts should be taken to increase the percentage of RA patients receiving DMARD treatment in accordance with guidelines.

Key messages
• One-fourth of German patients with RA are NP to MTX therapy.
• 6.5% of NP patients are also NA to MTX therapy.
• 50% of NP patients do not receive any follow-up DMARD treatment.

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Author contributions
Specifically, the main tasks the authors engaged in were TW, project lead, contributed to writing all parts of the article. SM contributed to statistical analysis and validation of database. UM/AF contributed to statistical analysis and interpretation of results in discussion section. J-PF, HH and KK contributed to conception/design of the study and interpretation of results in discussion section. All authors contributed toward data analysis, drafting and revising the paper and agree to be accountable for all aspects of the work.

Disclosure
TW has received honoraria from several pharmaceutical/consultancy companies (Novo Nordisk, Abbvie, Merck, GSK, BMS, LEO Pharma, Astra Zeneca, Bayer, Boehringer Ingelheim, Pharmerit). SM participated in this study as a staff member of IPAM; IPAM work in this study was sponsored by Roche Pharma AG and Chugai Pharma Europe Ltd, Germany. AF conducted consultancies for Bayer Pharma AG in 2016 and is employed by AOK PLUS. UM does not have any conflicts of interest except those potentially related to his employer, AOK PLUS. KK has received honoraria as a member of advisory boards and/or speakers bureau from: Abbvie, BMS, Medac, MSD, Pfizer, Roche and UCB. J-PF and HH are employed by Roche Pharma AG and Chugai Pharma Europe Ltd, German branch. The authors report no other conflicts of interest in this work.

References


17. Wilke T, Muller S, Morisky DE. Toward identifying the causes and combinations of causes increasing the risks of nonadherence to medical regimens: combined results of two German self-report surveys. Value Health. 2011;14(8):1092–1100.


Supplementary material

Figure S1 Distribution of MPR among RA patients who received MTX therapy.

Notes: The figure shows the distribution of patient-specific MPRs in the sample of RA patients who received MTX therapy. MPR was calculated during periods of therapy continuation only. So, our analysis was carried out from the first observed MTX prescription until the last observed MTX prescription. In case an MTX treatment gap was observed (>12 weeks), observation ended with the date of the last prescription before that observed gap.

Abbreviations: MPR, medication possession ratio; RA, rheumatoid arthritis; MTX, methotrexate.