Multidose drug dispensing and discrepancies between medication records

Liv Johanne Wekre, Olav Spigset, Olav Sletvold, Janne Kutschera Sund, Anders Grimsmo

INTRODUCTION
Incorrect use and incorrect handling of drugs is a major problem both in specialist care and in primary care. Even when limited to primary care, the number of adverse drug events has been seriously high in international studies. Both GPs and home-care services sense this problem and have expressed dissatisfaction with the collaboration regarding the medication of shared patients.

In 1999, the Norwegian Board of Health initiated work on a plan of action aimed towards safer use of medicines. The focus was especially directed towards primary care services. A number of actions was suggested, one of which was the use of MDD in nursing homes and home-care services. MDD implies that patients receive their drugs machine-dispensed into one unit for each dose occasion, packed into disposable bags. The dose unit bags are labelled with patient data, drug contents data and time for intake.

The number of MDD users is growing. In 2002, there were about 3000 MDD users in Norway; in 2006, the number had grown to 16 000 and in 2009 to approximately 35 000. Eighty per cent of the MDD users receive home-care service. There is an estimated potential of including a total of 200 000 patients from the primary care in Norway into the MDD system. MDD is expected to reduce medication errors, increase drug adherence and decrease waste of unused drugs. Still, more research is needed to document the effects of the system.

The aim of this study was to investigate whether the implementation of MDD for elderly outpatients was associated with a change in inconsistencies when comparing medication records from the GPs and the home-care services. We wanted to study both the number of discrepancies and the potential of the discrepancies to cause harm to the patients. To our knowledge, a controlled before–after study such as this has not been conducted before.

METHODS
Ten home care units in the city of Trondheim, Norway, each recruited up to 15 patients for participation, selecting the first 15 patients on an alphabetical list. The nurses responsible for the implementation of MDD in the units performed the selection and obtained informed consent. Medication records from the GPs and from the home-care services were collected half a year before and 1 year after the implementation of MDD. After the implementation, medication records from the pharmacies responsible for delivering the MD packages were provided as well. The study ran from May 2006 to January 2008.

The primary outcome was discrepancies between the patients’ medication records at the GPs and from the MD packages. The discrepancies were rated into four classes based upon the potential of including a total of 200 000 patients from the primary care in Norway into the MDD system. MDD is expected to reduce medication errors, increase drug adherence and decrease waste of unused drugs. Still, more research is needed to document the effects of the system.

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The primary outcome was discrepancies between the patients’ medication records at the GPs and from the home-care services. The discrepancies were rated for their potential to cause patient harm by a team consisting of two pharmacists, a GP, a clinical pharmacologist and a geriatrician. Each member of the team made an individual assessment.
before the joint evaluation. In cases of disagreement about the inconsistencies, the issue was resolved by discussion. Consensus was reached in all cases. The team members were blinded with regard to whether the records were collected before or after MDD was implemented.

Assessment was done by a validated method, and discrepancies were rated into one of three classes according to whether they had minimal, moderate or severe potential to harm. In addition, we included a fourth class of non-classifiable discrepancies (table 1).

Discrepancies caused both by discordant prescriptions and by missing information in the medication records were registered. Two approaches were used to study whether there were any significant changes before and after the implementation of MDD:

1. Comparing the sum of risk scores belonging to the pair of medication records from the GP and from the home-care services, before and after implementation of MDD. The sum of risk scores was calculated by giving a class 1 discrepancy 1 point, a class 2 discrepancy 2 points and a class 3 discrepancy 3 points. Class 0 discrepancies gave no points.

2. Comparing the number of high-risk medication records before and after implementation of MDD. High-risk medication records were defined by the following criteria:

   a. Records where the sum of risk-scores was 6 or higher.
   b. Records containing one or more class 3 discrepancies.

The drugs were divided into three groups: (1) drugs subject to MDD, (2) drugs not suitable for MDD and (3) drugs prescribed to be used as required. By doing this, we were able to set up an internal control. We could study whether changes in the number of discrepancies were present among drugs subject to MDD only or whether the changes observed were independent of the MDD per se.

Analyses were completed using Microsoft Office Excel 2003 (Microsoft, Seattle, Washington) and SPSS (version 16; SPSS, Chicago, Illinois) for Windows. The statistical analyses used were the Student t test for paired samples for continuous data and the McNemar test for paired nominal data. p Values <0.05 were considered statistically significant.

RESULTS

In total, 136 patients were included after the first collection of medication records. However, only 59 patients (43%) remained in the final material. The 77 drop-outs were as follows: 43 patients were not considered suitable for using MDD or did not receive home care when MDD was implemented, 20 patients had had MDD for a period of time but quit before the last collection of medication records, and 14 patients did receive MDD at the time of evaluation, but not all medication records were available (eight records missing from the GPs, three records missing from the home-care services and three records missing from the pharmacies). The patients had a mean age of 80 years at study start, ranging from 52 to 92 years. Forty-six (78%) of the patients were female. For comparison, the dropout patients had a mean age of 78 years, ranging from 54 to 98 years, and 56% were female.

The total number of drugs listed in the 59 medical records from the GPs was 386 before the implementation of MDD and 424 after the implementation (p = 0.016). Before the implementation, there were 47 medication records (80%) with discrepancies, as compared with 45 records (76%) with discrepancies after the implementation (p = 0.774). In total, there was a 34% reduction in the number of discrepancies after implementation of MDD (p < 0.001). The risk classification of the discrepancies is presented in table 2. For drugs subject to MDD, the reduction in the number of discrepancies was 59%, whereas for drugs not suitable for MDD (eg, injections, mixtures, eye-drops) and drugs to be used as required, the reduction was 31%. Table 3 shows the number of discrepancies for these three groups.

The various types of discrepancies are presented in table 4. The most frequent type of discrepancy both before and after implementation of MDD was that a prescription in the home-care services record was missing in the GP’s record. The second most frequent discrepancy was that a prescription in the GP’s record was lacking in the home-care services record. For all the 59 pairs of medication records, there was a total risk score of 308 before the implementation and 181 after the implementation of MDD (p < 0.001). There was also a significant

### Table 1 Classification of discrepancies according to potential harm

<table>
<thead>
<tr>
<th>Potential harm*</th>
<th>No of discrepancies before implementation, N = 386</th>
<th>No of discrepancies after implementation, N = 424</th>
<th>Absolute reduction in the percentage of discrepancies (95% CI)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Not classified (class 0)</td>
<td>12 (3%)</td>
<td>12 (3%)</td>
<td>0.29% (–1.15% to 1.73%)</td>
<td>0.690</td>
</tr>
<tr>
<td>Unlikely to cause discomfort (class 1)</td>
<td>84 (22%)</td>
<td>66 (15%)</td>
<td>6.66% (0.67% to 12.7%)</td>
<td>0.030</td>
</tr>
<tr>
<td>Potential to cause moderate discomfort (class 2)</td>
<td>97 (25%)</td>
<td>50 (12%)</td>
<td>12.3% (7.1% to 17.8%)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Potential to cause severe discomfort (class 3)</td>
<td>10 (3%)</td>
<td>5 (1%)</td>
<td>1.54% (–1.87% to 4.95%)</td>
<td>0.369</td>
</tr>
<tr>
<td>Total</td>
<td>203 (53%)</td>
<td>133 (31%)</td>
<td>21.0% (11.8% to 30.2%)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

*See text for a detailed explanation of the risk classification procedure. Number of prescriptions.
there were discrepancies in records belonging to patients
the implementation of MDD. A study from 2004 showed that
agreed with the GP record nor the home-care service record.
tions. In 31 drug prescriptions the pharmacy record neither
record and the home-care services record in 133 drug prescrip-
There were discrepancies between the pharmacy record and the
services were compared with the records from the pharmacies
information in the medication records from the GPs and at the
results from this study show a better agreement between the
a registered increase in the number of drugs in the medication
medication records at the home-care services compared with the
upto 90% of the patients had one or more discrepancies in their
medication records at the home-care services compared with the
Medication record at the GP,20 21 whereas we found 80% before
medication record at the GP,20 21 whereas we found that 76% of the
multidose patients had at least one discrepancy. A small study
from 2001 indicated that MDD causes no better agreement in
medication records than manual dispensing of drugs.8 This may
again indicate that the established routines in the home-care
services, at the GPs and at the pharmacies, together with the
information work accomplished during the implementation of
MDD, are important factors in order to achieve improved drug
safety in the MDD system.

The home-care services in Trondheim have used electronic
health records (EHRs) since 1996, and its medication module
regularly since 1998. This should be accounted for when
comparing the results with the findings from other studies24
since EHRs are not yet common everywhere.24 All medication
records from the community home-care services that we used in
the present study were printed out from the EHR.

Ninety-eight per cent of GPs in Norway have EHRs,25 but
even though all of them use the EHR when printing out single
prescriptions, the updating of the patient medication record has
not been carried out systematically. Before the implementation
of MDD, 10 of the included medication records from the GPs
were not printed out from the EHR medication module, as
compared with only one after the implementation. The better
routines among GPs in updating the medication records
probably contributed to the reduction in discrepancies that we
found.

We found a reduced number of discrepancies and a decrease in
estimated health risks due to discrepancies in the medication
records after the implementation of MDD. Studies show,
however, that a reduction in prescribing errors will not neces-
sarily be followed by a decrease in adverse drug events.26
Handling of the medication after removing it from the
packaging may still contribute to a high error frequency.27

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Number of discrepancies between the medication records from the general practitioner and from the home-care services before and after the implementation of multidose drug dispensing, classified on the basis of whether the drugs were dispensed in a multidose package or not</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type of discrepancy</td>
<td>No of discrepancies before implementation, N = 203</td>
</tr>
<tr>
<td>Multidose dispensable drugs</td>
<td>82 (21%)</td>
</tr>
<tr>
<td>Drugs not suitable for multidose dispensing</td>
<td>51 (13%)</td>
</tr>
<tr>
<td>Drugs to be used as required</td>
<td>70 (18%)</td>
</tr>
<tr>
<td>Total</td>
<td>203 (53%)</td>
</tr>
</tbody>
</table>

†Number of prescriptions.

A significant reduction in the number of discrepancies was
seen not only for the drugs subject to MDD, but also for the
drugs not suitable for MDD (table 3). This finding supports the
assumption that changes in routines constitute a central factor
in the improvement and that the improvement is not necessarily
due to the MDD alone. The improvement could be attributed to
the implementation process itself, and to the work done by the
different participants. The amount of allocated resources,
the mandate from the city council to the implementation team
and other routines adopted could also be of importance.

In previous Norwegian studies, it has been shown that up to
90% of the patients had one or more discrepancies in their
medication records at the home-care services compared with the
medication record at the GP,20 21 whereas we found 80% before
the implementation of MDD. A study from 2004 showed that
there were discrepancies in records belonging to patients
receiving MDD as well.23 Regarding the GPs’ and home-care
services’ records, discrepancies were disclosed in 52% of the
patients in that study,23 whereas we found that 76% of the
multidose patients had at least one discrepancy. A small study
from 2001 indicated that MDD causes no better agreement in
medication records than manual dispensing of drugs.8 This may
again indicate that the established routines in the home-care
services, at the GPs and at the pharmacies, together with the
information work accomplished during the implementation of
MDD, are important factors in order to achieve improved drug
safety in the MDD system.

DISCUSSION

Results from this study show a better agreement between the
information in the medication records from the GPs and at the
home-care services after the implementation of MDD compared
with before the implementation. This improvement caused a
drop in estimated health risks due to discrepancies in spite of
a registered increase in the number of drugs in the medication
records.

Table 4 | Type of discrepancies between the medication records from the general practitioner and from the home-care services before and after the implementation of multidose drug dispensing |
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Type of discrepancy</td>
<td>No of discrepancies before implementation, N = 203</td>
</tr>
<tr>
<td>Prescription lacking in the record from the general practitioner</td>
<td>83 (41%)</td>
</tr>
<tr>
<td>Prescription lacking in the record from the home-care services</td>
<td>0 (34%)</td>
</tr>
<tr>
<td>Different dosage</td>
<td>30 (15%)</td>
</tr>
<tr>
<td>Fixed prescription versus prescribed as required</td>
<td>7 (4%)</td>
</tr>
<tr>
<td>Different dose frequency*</td>
<td>3 (1%)</td>
</tr>
<tr>
<td>Missing information†</td>
<td>4 (2%)</td>
</tr>
<tr>
<td>Others</td>
<td>6 (3%)</td>
</tr>
<tr>
<td>Total</td>
<td>203 (100%)</td>
</tr>
</tbody>
</table>

*Different dose frequency but the same total daily dose, for example 50 mg × 2 versus 100 mg × 1.
†Missing information about type of formulation or drug dose in the prescription.
the other hand, the MDD system may reduce the number of prescribers, which is an independent risk factor. 28

Limitations of the study

The dropout rate was large, 57%. Since the study population was old, and not all patients are suited for MDD, this was expected. Twenty patients (15%) in the dropout group had received MDD in a period of time before the follow-up. We cannot exclude with certainty that some of them may have died from reasons connected to the implementation of MDD, but we do not have any information that this did happen.

Use of multiple statistical testing can inflate the type I error rate, so some of the statistically significant findings could be spurious, and the small sample size means that some possibly important differences could have been missed.

We considered the selection of patients from an alphabetical list to be convenient. As the selection was done by the last names of the patients, the risk of drawing family members could be increased. However, the patients were recruited from 10 different home care units, and we do not consider that this procedure has introduced any relevant bias.

The implementation process in Trondheim precluded the possibility of including a control of patients not subject to MDD from the same municipality. Including an external control group would involve a different organisation prone to be influenced by other factors in the study period. Instead, we made an internal control by comparing drugs subject to MDD and drugs not suitable for MDD. Changes in the latter group should not be caused by the introduction of the MDD system per se, but if such changes occurred, they should be due to other elements common to both groups.

Access to clinical data could have made the classification of the discrepancies more reliable. 17 18 However, the classification used in this study has also been used by others based on drug information data alone 16 and is validated for use in settings like the present one.

The GPs and the home care units were informed about the study, thus giving them the opportunity to scrutinise the lists before they were forwarded to the study investigators. Before the implementation of the MDD system, the home care personnel collected the medication records from the GPs and handed them over to the study investigators together with their own medication records. This procedure gave them an opportunity to change their own medication records, thereby omitting discrepancies. After the implementation of MDD, the study investigators contacted the GPs and the home-care services separately. The home-care services and the GPs could then not compare their records directly, but they had, at least in theory, the chance to check their records against the records they receive from the pharmacy when changes are made in the multidose packages. However, since the home care units were asked for several (until 15) medication records, a double check would be time-consuming. Hence, we consider such a scenario unlikely.

Finally, any generalisation of findings in a study from one single administration and organisation should be done with caution, and this is also the case for the findings in this study.

Acknowledgements

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Competing interests

None.

Patient consent

Obtained.

Ethics approval

Ethics approval was provided by the Regional Committee for Medical Research Ethics (REK) and the Norwegian Data Inspectorate (NSD).

Provenance and peer review

Not commissioned; externally peer reviewed.

REFERENCES

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