Monitoring of patients treated with lithium for bipolar disorder: an international survey

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Abstract

Background: Adequate monitoring of patients using lithium is needed for optimal dosing and for early identification of patients with (potential) ADEs. The objective was to internationally assess how health care professionals monitor patients treated with lithium for bipolar disorder.

Methods: Using networks of various professional organizations, an anonymous online survey was conducted among health care professionals prescribing lithium. Target lithium serum levels and frequency of monitoring was assessed together with monitoring of physical and laboratory parameters. Reasons to and not to monitor and use of guidelines and institutional protocols, and local monitoring systems were investigated.

Results: The survey was completed by 117 health care professionals incorporating responses from twenty-four countries. All prescribers reported to monitor lithium serum levels on a regular basis, with varying target ranges. Almost all (> 97%) monitored thyroid and renal function before start and during maintenance treatment. Reported monitoring of other laboratory and physical parameters was variable. The majority of respondents (74%) used guidelines or institutional protocols for monitoring. In general, the prescriber was responsible for monitoring, had to request every monitoring parameter separately and only a minority of patients was automatically invited.

Conclusions: Lithium serum levels, renal and thyroid function were monitored by (almost) all physicians. However, there was considerable variation in other monitoring parameters. Our results help to understand why prescribers of lithium monitor patients and what their main reasons are not to monitor patients using lithium.

Keywords: Lithium, Bipolar disorder, Survey, Monitoring, Therapeutic drug monitoring

Background

Mood stabilizers play a pivotal role in the long-term treatment of patients with bipolar disorder. While it is known that mood stabilizers effectively treat symptoms of bipolar disorder, the potential for adverse drug events (ADEs) is of concern (Rothschild et al. 2007; Ayani et al. 2016; Mann et al. 2008). Some ADEs are preventable, and steps to minimize errors are needed to improve patient safety (Leendertse et al. 2008).

Lithium is the gold standard for maintenance treatment of patients with bipolar disorder (Sani et al. 2017). It is a drug with a narrow therapeutic index; therefore, careful therapeutic drug monitoring is needed to maximize effectiveness and to minimize ADEs and toxicity. Patients using lithium are known to have a high intra- and inter-patient variability in dose—concentration relationship, and external factors including drug–drug interactions, environmental temperature, and fluid and electrolyte intake may influence lithium serum levels (Amdisen 1980; Huang et al. 2008; Wilting et al. 2005; Wilting et al. 2007; Rej et al. 2014). Recommendations regarding optimal lithium serum levels differ among clinical practice guidelines, with the most common range being between 0.6 and 0.8 mmol/L for maintenance treatment (Malhi...
et al. 2017). The illness stage may further require a different approach for lithium serum levels (Malhi et al. 2016).

Besides monitoring of lithium serum levels, clinical and biomarker monitoring of physical and laboratory parameters is required (Nederlof et al. 2015). Lithium use has been associated with a gradual decline in renal function: estimated glomerular filtration rate (eGFR) decreases by about 30% more than associated with aging alone (Tondo et al. 2017). This decline in renal function can become irreversible and may lead to renal failure (Gitlin 2016). Moreover, a decline in eGFR can lead to an increased risk of lithium toxicity due to accumulation (Lepkifker et al. 2004). Furthermore, lithium is known to affect thyroid function (van Melick et al. 2010); with up to 40% of patients experiencing goiter and about 20% hypothyroidism (Lazarus 2009), although the reported prevalence shows a wide range. A 10% higher value of parathyroid hormone (PTH) and calcium was found in a meta-analysis comparing patients using lithium to healthy controls or psychiatric patients not treated with lithium (McKnight et al. 2012). These elevations could stimulate pathological changes leading to the occurrence of hyperplasia or parathyroid adenomas (Giusti et al. 2012). In addition, physical parameters such as weight and blood pressure may be influenced during lithium use (Gitlin 2016; Bisogni et al. 2016).

Adequate monitoring of patients during lithium use is needed for early identification of patients with (potential) ADEs and for optimal dosing (Kirkham et al. 2013; Severus et al. 2008; Aral and Vecchio-Sadus 2008; Baird-Gunning et al. 2017). So far, it is not known how different health care professionals internationally aim to achieve safe and effective lithium treatment through monitoring. Local monitoring systems may differ and health care professionals may be influenced by different guidelines, institutional or laboratory protocols, scientific literature, and their own personal knowledge and experience. The objective of this study was therefore to internationally assess how health care professionals report to monitor patients treated with lithium.

**Methods**

**Study design, participants and recruitment**

An international survey was conducted between October 2016 and April 2017 among health care professionals prescribing lithium to patients with bipolar disorder. Health care professionals were recruited through networks of various professional organizations (Additional file 1: Appendix S1), and invited by email to participate. The email introduced the nature and purpose of the survey and proposed to further forward the survey to lithium prescribing colleagues. To maximize the response rate, attention was generated during presentations for the Dutch Foundation for Bipolar Disorders and the Belgian College of Neurological and Biological Psychiatry and through flyers at a meeting of the International Group for The Study of Lithium Treated Patients (IGSLi). The survey was conducted in English. Participation was voluntary and answers were kept with anonymized identity of participants. A reminder email was sent a few weeks following the initial request. Study data were analyzed at the Division of Pharmacoepidemiology and Clinical Pharmacology of Utrecht University. The study protocol was approved by the Institutional Review Board of the Division of Pharmacoepidemiology and Clinical Pharmacology of Utrecht University.

**Questionnaire**

Based on monitoring recommendations in various guidelines, a questionnaire was designed to assess how health care professionals prescribing lithium to patients with bipolar disorder report to monitor these patients. The primary researcher (MN) assessed all monitoring instructions in four guidelines (Kupka et al. 2015; National Institute for Health and Care Excellence 2014; Ng et al. 2009; Grunze et al. 2013), to explore relevant monitoring parameters and advised frequencies of monitoring: the Dutch Multidisciplinary guideline (Kupka et al. 2015), the International Society for Bipolar Disorders guidelines (Ng et al. 2009) and the World Federation of Societies of Biological Psychiatry guidelines (Grunze et al. 2013), and the guideline of the National Institute for Health and Care Excellence (2014). Actual utilization of guidelines was assessed, including explicationary reasons. In addition, we included questions on local monitoring systems. Most questions were multiple choice with an option offered for comments at the end, if relevant. The study group then composed a questionnaire in LimeSurvey version 2.5, an online accessible survey application. The survey was tested by eleven Dutch experts in the field of lithium treatment (Additional file 2: Appendix S2) to optimize content on appropriateness, relevance, and time to complete the survey, and it was adjusted accordingly.

The final questionnaire consisted of 41 questions divided into four parts. First, target lithium serum levels and frequency of monitoring were assessed. Second, monitoring frequency of physical and laboratory parameters was assessed. Third, to identify factors contributing to monitoring, reasons to monitor or not to monitor, and use of guidelines or institutional protocols were investigated. Fourth, personal characteristics of the respondent including age, gender, experience with prescribing lithium, and country of residence were collected. A condensed version of the questionnaire can be found in...
Table 1. The full questionnaire can be found in Additional file 3: Appendix 3.

Analysis

Responses were analyzed using the Statistical Package for Social Sciences Version 24.0 for Windows (SPSS 24.0; SPSS Inc, Chicago, IL). Data were analyzed using descriptive statistics. Target lithium serum levels were presented with the use of boxplot statistics using Tukey’s Hinges (Tukey 1977).

Results

Study sample

A total of 117 respondents completed the survey. The majority of respondents were employed as psychiatrists (91%), followed by nurse practitioners (3%), psychiatric residents (5%), or as general practitioner (1%). Twenty percent of the respondents were self-employed. The duration of experience with prescribing lithium had a median of 22 years [interquartile range (IQR) 10–33]. Two-third (66%) of the respondents were male; the median age was 53 years (IQR 41–60).

Responses were collected from health care professionals residing in twenty-four different countries. Countries where prescribers were employed were diverse, with 63% of the respondents employed in Europe, 22% in North America, 9% in South America, 3% in Australia, 2% in Asia and 1% in Africa (Fig. 1).

Lithium serum level and frequency of monitoring

In Fig. 2, target values for minimum and maximum lithium plasma levels the respondents reported to pursue are presented for acute and maintenance treatment in adults and elderly patients.

For adults, the median lower target level for acute treatment was 0.8 mmol/L (IQR 0.1; range 0.5–1.0). The median for the upper level was 1.1 mmol/L (IQR 0.2; range 0.7–1.6). For maintenance treatment the median target serum level for adults was 0.6 mmol/L (IQR 0.1; range 0.4–0.8); with a median upper level of 0.8 mmol/L (IQR 0.1; range 0.5–1.2).

For elderly, the median target range for serum levels was 0.6 mmol/L (IQR 0.2; range 0.3–1.0), with 0.9 mmol/L (IQR 0.2; range 0.5–1.2) as upper level for acute treatment. During maintenance treatment the median lower level was 0.4 mmol/L (IQR 0.1; range 0.3–0.8); with 0.8 mmol/L (IQR 0.2; range 0.6–1.2) as upper level. Spread in target ranges for both adults and elderly patients during acute treatment was relatively larger compared to maintenance treatment.

All respondents monitored lithium serum levels during treatment, although there was variability in frequency. In the initial phase of lithium treatment, most respondents monitored lithium serum levels one to three times in the first month and between one and three times during months 2–6. During maintenance treatment half of the respondents monitored one to three times a year; others four or more times a year (44%) (Table 2). Many had difficulties explaining how often they monitored the lithium serum level during the first month, as they monitored lithium serum levels until they reached the target level. Thirteen respondents commented to monitor 5–7 days after every dose change.
Monitoring of physical and laboratory parameters

All responders monitored at least one physical or laboratory parameter. Responses of monitoring of physical and laboratory monitoring are shown in Table 3.

Most respondents monitored bodyweight, blood pressure, pulse, and body mass index. Fewer respondents
### Table 3 Monitoring of physical and laboratory parameters

| Parameter                              | Monitoring before start of treatment | Frequency of monitoring during the first 6 months of treatment | Frequency of monitoring per year during maintenance treatment |
|----------------------------------------|--------------------------------------|---------------------------------------------------------------|---------------------------------------------------------------|
|                                        | Never | 1–3 x | ≥ 4 x | Never | Once | 2–3 x | ≥ 4 x |
| Physical parameters                    |       |       |       |       |       |       |       |
| Physical parameters (at least one parameter) | 104 (88%) | 17 (14%) | 76 (64%) | 24 (21%) | 12 (10%) | 58 (49%) | 28 (24%) | 19 (16%) |
| Body weight                            | 92 (79%) | 32 (27%) | 70 (60%) | 15 (13%) | 20 (17%) | 58 (50%) | 25 (21%) | 14 (12%) |
| Blood pressure                         | 88 (75%) | 35 (30%) | 63 (54%) | 19 (16%) | 28 (24%) | 51 (44%) | 24 (21%) | 14 (12%) |
| Pulse                                  | 79 (68%) | 43 (37%) | 56 (48%) | 18 (15%) | 38 (32%) | 48 (41%) | 19 (16%) | 12 (10%) |
| Body mass index (BMI)                  | 63 (54%) | 54 (46%) | 54 (46%) | 9 (8%) | 45 (38%) | 47 (40%) | 16 (14%) | 9 (8%) |
| Electrocardiogram (ECG)                | 51 (44%) | 66 (56%) | 46 (39%) | 5 (4%) | 71 (61%) | 31 (27%) | 12 (10%) | 3 (3%) |
| Waist circumference                    | 45 (38%) | 72 (62%) | 42 (36%) | 3 (3%) | 65 (56%) | 38 (32%) | 9 (8%) | 5 (4%) |
| Pregnancy test                         | 40 (34%) | 90 (77%) | 24 (21%) | 3 (3%) | 102 (87%) | 7 (6%) | 6 (5%) | 2 (2%) |
| 24 h urine examination                 | 23 (20%) | 92 (79%) | 21 (18%) | 3 (3%) | 88 (75%) | 23 (20%) | 2 (2%) | 4 (3%) |
| Renal function                         |       |       |       |       |       |       |       |       |
| Renal function (at least one parameter) | 116 (99%) | 3 (3%) | 88 (75%) | 26 (22%) | 0 | 25 (21%) | 60 (51%) | 32 (27%) |
| Creatinine                             | 116 (99%) | 3 (3%) | 88 (75%) | 26 (22%) | 0 | 26 (22%) | 59 (50%) | 32 (27%) |
| Urea                                   | 101 (86%) | 21 (18%) | 78 (67%) | 18 (15%) | 18 (15%) | 22 (19%) | 48 (41%) | 29 (25%) |
| Glomerular filtration rate (GFR)       | 100 (85%) | 19 (16%) | 76 (65%) | 22 (19%) | 13 (11%) | 22 (19%) | 52 (44%) | 30 (26%) |
| Albumin                                | 66 (56%) | 46 (39%) | 60 (51%) | 11 (9%) | 42 (36%) | 21 (18%) | 34 (29%) | 20 (17%) |
| Thyroid function                       |       |       |       |       |       |       |       |       |
| Thyroid function (at least one parameter) | 115 (98%) | 2 (2%) | 104 (89%) | 11 (9%) | 0 | 43 (36%) | 54 (46%) | 20 (17%) |
| Thyroid stimulating hormone (TSH)      | 115 (98%) | 2 (2%) | 104 (89%) | 11 (9%) | 1 (1%) | 42 (36%) | 54 (46%) | 20 (17%) |
| Thyroxine (T4)                         | 82 (70%) | 30 (26%) | 80 (68%) | 7 (6%) | 28 (24%) | 33 (28%) | 41 (35%) | 15 (13%) |
| Parathyroid function                   |       |       |       |       |       |       |       |       |
| Parathyroid hormone (PTH)              | 32 (27%) | 79 (68%) | 35 (30%) | 3 (3%) | 72 (62%) | 22 (19%) | 17 (15%) | 6 (5%) |
| Electrolytes                           |       |       |       |       |       |       |       |       |
| Electrolytes (at least one parameter)  | 107 (91%) | 10 (9%) | 86 (73%) | 21 (18%) | 8 (7%) | 34 (29%) | 49 (42%) | 26 (22%) |
| Sodium                                 | 103 (88%) | 17 (15%) | 80 (68%) | 20 (17%) | 14 (12%) | 32 (27%) | 46 (39%) | 25 (21%) |
| Potassium                              | 101 (86%) | 18 (15%) | 79 (68%) | 20 (17%) | 15 (13%) | 32 (27%) | 45 (38%) | 25 (21%) |
| Calcium                                | 94 (80%) | 21 (18%) | 81 (69%) | 15 (13%) | 19 (16%) | 35 (30%) | 44 (38%) | 19 (16%) |
| Hematological parameters               |       |       |       |       |       |       |       |       |
| Hematological parameters (at least one parameter) | 106 (90%) | 26 (22%) | 79 (67%) | 12 (10%) | 24 (21%) | 51 (44%) | 29 (25%) | 13 (11%) |
| Full blood count                       | 102 (87%) | 32 (27%) | 73 (62%) | 12 (10%) | 30 (26%) | 47 (40%) | 29 (25%) | 11 (9%) |
| Leukocytes                             | 101 (86%) | 30 (26%) | 75 (64%) | 12 (10%) | 27 (23%) | 50 (43%) | 27 (23%) | 13 (11%) |
| Leukocyte differentiation              | 87 (74%) | 40 (34%) | 67 (57%) | 10 (9%) | 38 (32%) | 45 (38%) | 25 (21%) | 9 (8%) |
| Lipid profile                          |       |       |       |       |       |       |       |       |
| Lipid profile (at least one parameter) | 73 (62%) | 57 (49%) | 55 (47%) | 5 (4%) | 42 (36%) | 53 (45%) | 18 (15%) | 4 (3%) |
| Triglycerides                          | 71 (61%) | 58 (50%) | 54 (46%) | 5 (4%) | 43 (37%) | 52 (44%) | 18 (15%) | 4 (3%) |
| Total cholesterol                      | 71 (61%) | 59 (50%) | 53 (45%) | 5 (4%) | 44 (38%) | 52 (44%) | 17 (15%) | 4 (3%) |
| High density lipoprotein (HDL)         | 69 (59%) | 60 (51%) | 53 (45%) | 4 (3%) | 45 (38%) | 51 (44%) | 18 (15%) | 3 (3%) |
| Low density lipoprotein (LDL)          | 67 (57%) | 62 (54%) | 50 (43%) | 4 (3%) | 48 (41%) | 49 (42%) | 17 (15%) | 3 (3%) |
| Very low density lipoprotein (vLDL)    | 35 (30%) | 84 (72%) | 32 (27%) | 1 (1%) | 71 (61%) | 35 (30%) | 11 (9%) | 0 |
| Hepatic function                       |       |       |       |       |       |       |       |       |
| Hepatic function (at least one parameter) | 68 (58%) | 61 (52%) | 48 (41%) | 8 (7%) | 54 (46%) | 37 (32%) | 20 (17%) | 6 (5%) |
| Alanine aminotransferase (ALAT)        | 68 (58%) | 62 (53%) | 47 (40%) | 8 (7%) | 55 (47%) | 37 (32%) | 19 (16%) | 6 (5%) |
| Aspartate aminotransferase (ASAT)      | 66 (56%) | 64 (53%) | 46 (39%) | 7 (6%) | 55 (47%) | 37 (32%) | 19 (16%) | 6 (5%) |
| Bilirubin                              | 34 (29%) | 79 (68%) | 33 (28%) | 5 (4%) | 71 (61%) | 27 (23%) | 15 (13%) | 3 (3%) |
| Other laboratory parameters            |       |       |       |       |       |       |       |       |
| Fasting glucose                        | 79 (67%) | 48 (41%) | 63 (54%) | 6 (5%) | 35 (30%) | 56 (48%) | 20 (17%) | 6 (5%) |

Italicized cells represent most reported answers after initiation of lithium treatment.
made electrocardiograms, monitored waist circumference, performed pregnancy tests, or 24 h urine examinations. Renal function and thyroid function were monitored by almost all respondents before start and during maintenance treatment. Primarily creatinine, thyroid stimulating hormone (TSH), urea, and glomerular filtration rate were measured. PTH was considerably less often monitored, sometimes solely in case of a deviating calcium level. A few mentioned to measure anti-thyroid autoantibodies (n = 6), mostly in case of abnormal TSH. Electrolytes and hematological parameters were generally determined before start and often repeated during maintenance treatment. Slightly more than half of the respondents monitored a lipid profile and hepatic function before start. Most of them never monitored hepatic function after start, while lipids were monitoring more often once a year during maintenance treatment. Fasting glucose was monitored by about two-thirds of the respondents before start, and about half of the respondents monitored fasting glucose one to three times during the first 6 months and thereafter once a year during maintenance treatment.

Some respondents mentioned to monitor the following additional parameters: alkaline phosphatase (ALP) (n = 1), CRP (n = 1), cysteine (n = 1), EEG (n = 1), erythrocyte sedimentation rate (n = 1), folate (n = 2), gamma-glutamyltransferase (GGT) (n = 1), glycated hemoglobin (HbA1c) (n = 1), serum level of antipsychotic agents (n = 1), urine osmolality (n = 1) vitamin D (n = 4), physical examination (n = 1), vitamin B1 or vitamin B12 (n = 2).

**Reasons for monitoring**
The main reason to monitor patients using lithium was because of safety or adverse effects (99%). Respondents monitored additionally to aid in dose adjustment or optimization (94%), in case of possible lithium toxicity (85%), during start, stop, or dose-adjustment of interacting medication (67%), to verify therapeutic level, especially in case of insufficient efficacy (64%), due to co-morbidities of the patient (63%), and because it is recommended in guidelines (59%). Other mentioned reasons that were to assess compliance (3%), to monitor patients during pregnancy (1%) or in case of a new episode (1%), or if a medication brand changed (1%).

Reasons not to monitor patients during lithium treatment were a lack of resources to monitor (4%), non-cooperative patients (4%), if a guideline did not oblige monitoring (3%), if monitoring was unnecessary in the opinion of the respondent (3%), if an institutional protocol did not require monitoring (2%), or unawareness of the necessity of monitoring (2%). Still, 90% of the respondents reported they always monitored their patients. One psychiatrist commented that there are no good reasons not to monitor a patient on lithium.

**Use of guidelines or institutional protocols for monitoring**
The monitoring policy of 74% of the respondents was based upon a guideline or institutional protocol. A minority of respondents said not to use guidelines because personal experience and practice were more valuable to them (9%). A number of respondents thought guidelines were not explicit enough (7%) or the institutional protocol or Summary of Product Characteristics (SmPC) they used was sufficient. Alternative reasons not to use guidelines were difficulties to apply guidelines to their patient population (3%) or an unawareness of the existence of such guidelines (2%). Additionally, respondents mentioned to perform monitoring based on individual patient characteristics (2%) or to experience a difficulty to choose between guidelines (1%). Sometimes guidelines were found to be excessive (1%) or they used their own reading of literature (1%).

**Local systems for ensuring monitoring patients treated with lithium**
The majority of respondents reported to be personally responsible for monitoring (91%). The health care professional responsible for monitoring was a psychiatrist (83%), psychiatric resident (3%), nurse practitioner (3%), or the head/clinic nurse (3%). A shared responsibility was mentioned by some respondents (3%). In a few responses, a general practitioner (2%), family physician (1%) or the doctor who continued treatment after discharge (1%) was responsible for monitoring.

Among the respondents, 74% had to request each parameter separately. For monitoring a combination of parameters during lithium treatment, 25% could make use of an existing laboratory protocol. For some health care professionals, the system for ensuring monitoring was organized in a way that patients were automatically invited by a laboratory or physician for determination of monitoring parameters (26%).

**Discussion**
We surveyed prescribers of lithium on monitoring of patients using lithium for bipolar disorder. All prescribers reported to monitor lithium serum levels on a regular basis, with varying intervals and target levels. Target levels for maintenance treatment were mostly consistent with target ranges in guidelines for bipolar disorders (Malhi et al. 2017). Target levels for acute treatment were...
more variable, which is in accordance with the variability among guidelines (Malhi et al. 2017).

Almost all health care professionals monitored thyroid and renal function before start, and all during maintenance treatment, indicating a higher priority was given to these parameters compared to other parameters. This does not mean that all respondents gave less priority to other monitoring physical or laboratory parameters. Some may consider other parameters equally important, but our results show that they reported to monitor at least renal and thyroid function. Respondents may not monitor these parameters in all patients using lithium in clinical practice. From studies in The Netherlands and the UK it is known that monitoring rates may be much lower than the health care professionals reported in our survey; showing that 8–22% of patients did not have any recorded test for renal or thyroid function (van de Beek et al. 2010; Paton et al. 2013). A difference between reported and actual monitoring in clinical practice can be a consequence of inadequate adherence to monitoring by patients (Collins et al. 2010). It is possible that respondents despite anonymity may have given socially desirable answers to our survey. If patients using lithium are not adequately monitored, ADEs may not be identified in time. On the other hand, excessive monitoring may have a negative impact on health care costs and provide an extra burden to patients and may even lead to a reduced treatment adherence (Tran et al. 2015).

Reported monitoring was more variable for other physical and laboratory parameters than thyroid and renal function. It would be of interest to assess reasons of health care professionals to monitor or not to monitor these parameters. Variability in monitoring between respondents can be a consequence of a lack of resources, difference in education, type of patient population, a different opinion on the need for monitoring or differences in recommendations in guidelines (Zivanovic 2017). The number of responses was not sufficient to compare responses between subgroups such as countries, type of health care professional or to determine guideline adherence.

Overall, respondents reported to use guidelines or institutional protocols for monitoring. Guidelines differ with respect to their recommendations, and studies have shown that there are discrepancies between guideline recommendations and actual monitoring in clinical practice (Malhi et al. 2017; van de Beek et al. 2010; Paton et al. 2013; Minay et al. 2013). Reasons to use or not to use guidelines were assessed in our survey. In line with previous research, several respondents reported not to follow guidelines because in their opinion they did not apply to their patient population (Perlis 2007). Guidelines are mainly based on literature such as randomized controlled trials that may not include specific patient populations, for instance children or older patients (Zarin et al. 2005). In our survey, health care professionals mentioned personal experience and practice to be the main reasons for not following a guideline. A lack of resources and non-cooperative patients could further complicate monitoring. Not all laboratories are equipped to perform all laboratory tests (Guo et al. 2013) and for prescribers of lithium not all monitoring facilities may be easily accessible, for example for making an electrocardiogram.

In the majority of cases the prescriber of lithium, generally a psychiatrist, was responsible for monitoring. The health care system should be organized in a way that continuity of monitoring is warranted. For instance, in the Dutch guideline, the prescriber is appointed to be responsible for monitoring (Kupka et al. 2015), but another health care professional can have a coordinating function. Among respondents, a small group reported that patients were automatically invited by a laboratory or physician for monitoring. Most respondents had to request each monitoring parameter separately, which may increase the workload and demands awareness of health care professionals. A method to order all monitoring parameters after a prescription has been described, known as a corollary order (Wright and Sittig 2006). In this way, an order set of monitoring parameters could be developed to be used when a health care professional prescribes lithium. Clinical decision systems could be programmed in a way that all relevant monitoring parameters are automatically ordered after a lithium prescription.

Strengths and limitations
An advantage of using a survey is that we could assess anonymized reported monitoring and use of guidelines. Our results help to understand why prescribers of lithium monitor patients and what their main reasons are not to monitor patients according to guideline recommendations or even not at all. By using an international online survey we were able to include respondents from 24 countries worldwide.

A limitation is that the response was not sufficient to compare individual countries. The limited response could have been a result of a limited amount of time or a lack of interest in the subject. Since the survey was distributed via various networks, we do not know how many professionals actually have been reached. Prescribers who did participate in the survey, may have been a selection of prescribers who are more interested in the subject. The survey was sent to health care professionals who were members of professional health care organizations,
including those with a specific focus on bipolar disorder. It is expected that they are better informed on lithium monitoring as some of them may have a special interest in lithium treatment. Respondents were relatively experienced, with a median of 22 years of prescribing lithium. Therefore, responses may not reflect monitoring in general clinical practice internationally.

Conclusions
Adequate monitoring of patients using lithium is needed for optimal dosing and early identification of patients with (potential) ADEs. In our survey, health care professionals prescribing lithium report to monitor lithium serum levels, renal and thyroid function on a regular basis. However, there was considerable variation in other monitoring parameters. Our results help to understand why prescribers of lithium monitor patients and what their main reasons are not to monitor patients. In future, it will be of interest to assess how prescribers respond to deviating parameters.

Additional files

Additional file 1: Appendix S1. Networks of professional organizations.
Additional file 2: Appendix S2. List of Dutch lithium experts who contributed to the pilot study.
Additional file 3: Appendix S3. Lithium survey.

Abbreviations
ADE: adverse drug event; ALP: alkaline phosphatase; eGFR: estimated glomerular filtration rate; GGT: gamma-glutamyltransferase; HbA1c: glycated hemoglobin; IGLLS: International Group for The Study of Lithium Treated Patients; IQR: interquartile range; PTH: parathyroid hormone; SmPC: Summary of Product Characteristics; TSH: thyroid stimulating hormone.

Authors’ contributions
MN is the primary author and performed the survey, analyzing, writing and editing of the data. RK, EH, TE, IW, LS and RH helped spreading the survey and provided critical appraisal of the manuscript. All authors read and approved the final manuscript.

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Competing interests
The authors declare that they have no competing interests.

Availability of data and materials
The original data is available at the Division of Pharmacoepidemiology and Clinical Pharmacology, Utrecht Institute for Pharmaceutical Sciences, Utrecht University, Utrecht, The Netherlands.

Consent for publication
Not applicable.

Ethics approval and consent to participate
The study protocol was approved by the Institutional Review Board of the Division of Pharmacoepidemiology and Clinical Pharmacology of Utrecht University.

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References

Amdisen A. Serum concentration and clinical supervision in monitoring of lithium treatment. Ther Drug Monit. 1980;2(1):73–83.

Arnal H, Vecchio-Sadus A. Toxicity of lithium to humans and the environment—a literature review. Ecotoxicol Environ Saf. 2008;70(3):349–56.

Ayani N, Sakuma M, Morimoto T, Kikuchi T, Watanabe K, Narumoto J, et al. The epidemiology of adverse drug events and medication errors among psychiatric inpatients in Japan: the JADE study. BMC Psychiatry. 2016;30(16):303.

Baird-Gunning J, Lea-Henry T, Hoegberg LCG, Gosselin S, Roberts DM. Lithium poisoning. J Intensive Care Med. 2017;32(4):249–63.

Bisognin V, Rossetto G, Reghin F, Padrini R, Rossi GP. Antihypertensive therapy in patients on chronic lithium treatment for bipolar disorders. J Hypertens. 2016;34(1):20–8.

Collins N, Barnes TR, Shingleton-Smith A, Gerrett D, Paton C. Standards of lithium monitoring in mental health trusts in the UK. BMC Psychiatry. 2010;12(10):80.

Gitin M. Lithium side effects and toxicity: prevalence and management strategies. Int J Bipolar Disord. 2016;4(1):27.

Giusti CF, Amorim SR, Guerra RA, Portes ES. Endocrine disturbances related to the use of lithium. Arq Bras Endocrinol Metabol. 2012;56(3):153–8.

Grunze H, Vleka E, Goodwin GM, Bowden C, Litch RW, Möller H-J, et al. The World Federation of Societies of Biological Psychiatry (WFSBP) guidelines for the biological treatment of bipolar disorders: update 2012 on the long-term treatment of bipolar disorder. World J Biol Psychiatry Off J World Fed Soc Biol Psychiatry. 2013;14(3):154–219.

Guo W, Guo G-X, Sun C, Zhang J, Rong Z, He J, et al. Therapeutic drug monitoring of psychotropic drugs in China: a nationwide survey. Ther Drug Monit. 2013;35(6):816–22.

Huang HC, Chang YT, Lin TH, Chiu HJ, Liu WM, Lee TJF. Prediction of optimal lithium doses for Taiwanese psychiatric patients. J Clin Pharm Ther. 2008;33(2):115–21.

Kirkham E, Bazire S, Anderson T, Wood J, Grassby P, Desborough JA. Impact of active monitoring on lithium management in Norfolk. Ther Adv Psychopharmacol. 2013;3(5):260–5.

Kupka RW, Goossens P, van Bendegem M, Daemen P, Daggenvoorde T, Daniels M, Dols A, van Dun D, Hillesges M, Hoogelande A, ter Kuive E, Pheottom T, Schulte R, Stevens A. Dutch multidisciplinary guideline bipolar disorder. Nederlandse Vereniging voor Psychiatrie (NVvP). 2015. http://www.nvvp.net/stream/richtlijn-bipolaire-stoornissen-2015. Accessed 15 Sept 2016.

Lazarus JH. Lithium and thyroid. Best Pract Res Clin Endocrinol Metab. 2009;23(6):723–33.
Leendertse AJ, Egberts ACG, Stoker LJ, van den Bemt PMLA, HARM Study Group. Frequency of and risk factors for preventable medication-related hospital admissions in the Netherlands. Arch Intern Med. 2008;168(17):1890–6.

Lepkifker E, Sverdlik A, Iancu I, Ziv R, Segev S, Koterl M. Renal insufficiency in long-term lithium treatment. J Clin Psychiatry. 2004;65(6):850–6.

Malhi GS, Gershon S, Outhred T. Lithiummeter: version 2.0. Bipolar Disord. 2016;18(8):631–41.

Malhi GS, Gershon S, Outhred T. The use of lithium for the treatment of bipolar disorder: recommendations from clinical practice guidelines. J Affect Disord. 2017;227(17):266–80.

Mann K, Rothschild JM, Keohane CA, Chu JA, Bates DW. Adverse drug events and medication errors in psychiatry: methodological issues regarding identification and classification. World J Biol Psychiatry Off J World Fed Soc Biol Psychiatry. 2008;9(1):24–33.

McKnight RF, Adida M, Budge K, Stockton S, Goodwin GM, Geddes JR. Lithium toxicity profile: a systematic review and meta-analysis. Lancet Lond Engl. 2012;379(9817):721–8.

Minay J, Paul R, McGarvey D, Savage G, Stevenson M, Fogarty D, et al. Lithium usage and renal function testing in a large UK community population; a case-control study. Gen Hosp Psychiatry. 2013;35(6):631–5.

National Institute for Health and Care Excellence. NICE guideline bipolar disorder; 2014. http://www.nice.org.uk/guidance/cg185/. Accessed 15 Sept 2016.

Nederlof M, Stoker LJ, Egberts TCG. Instructions for clinical and biomarker monitoring in the summary of product characteristics (SmPC) for psychotropic drugs: overview and applicability in clinical practice. J Psychopharmacol Oxf Engl. 2015;29(12):1248–54.

Ng F, Mammon OK, Wilting J, Sachs GS, Ferrier IN, Cassidy F, et al. The International Society for Bipolar Disorders (ISBD) consensus guidelines for the safety monitoring of bipolar disorder treatments. Bipolar Disord. 2009;11(6):559–95.

Paton C, Adroer R, Barnes TRE. Monitoring lithium therapy: the impact of a quality improvement programme in the UK. Bipolar Disord. 2013;15(8):865–73.

Perlis RH. Use of treatment guidelines in clinical decision making in bipolar disorder: a pilot survey of clinicians. Curr Med Res Opin. 2007;23(3):467–75.

Rej S, Alageel B, Segal M, Low NCP, Mucsi I, Holcroft C, et al. Is environmental temperature related to renal symptoms, serum lithium levels, and other laboratory test results in current lithium users? Hum Psychopharmacol. 2014;29(4):392–6.

Rothschild JM, Mann K, Keohane CA, Williams DH, Foskett C, Rosen SL, et al. Medication safety in a psychiatric hospital. Gen Hosp Psychiatry. 2007;29(2):156–62.

Sani G, Perugi G, Tondo L. Treatment of bipolar disorder in a lifetime perspective: is lithium still the best choice? Clin Drug Invest. 2017;37(8):13–27.

Severus WE, Kleindienst N, Seemüller F, Frangou S, Möller HJ, Greil W. What is the optimal serum lithium level in the long-term treatment of bipolar disorder—a review? Bipolar Disord. 2008;10(2):231–7.

Tondo L, Abramowicz M, Alda M, Bauer M, Bocchetta A, Bolzani L, et al. Long-term lithium treatment in bipolar disorder: effects on glomerular filtration rate and other metabolic parameters. Int J Bipolar Disord. 2017;5(1):27.

Tran VT, Barnes C, Montori VM, Falissard B, Ravaud P. Taxonomy of the burden of treatment: a multi-country web-based qualitative study of patients with chronic conditions. BMC Med. 2015;14(13):115.

Tukey JW. Exploratory data analysis. Boston: Addison-Wesley; 1977.

van de Beek LM, Ouwens MA, de Vries PL, Hummelen JW. Lithium levels need to be monitored: discrepancies between guidelines and practice. Tijdschr Voor Psychiatr. 2010;52(6):367–73.

van Melick EJM, Wilting J, Meinders AE, Egberts TCG. Prevalence and determinants of thyroid disorders in elderly patients with affective disorders: lithium and nonlithium patients. Am J Geriatr Psychiatry Off J Am Assoc Geriatr Psychiatry. 2010;18(5):395–403.

Wilting J, Movig KL, Zoelenaar MA, Hekster YA, Brouwers JR, Heerdink ER, et al. Drug-drug interactions as a determinant of elevated lithium serum levels in daily clinical practice. Bipolar Disord. 2005;7(3):274–80.

Wilting J, Fase S, Martens EP, Heerdink ER, Nolen WA, Egberts ACG. The impact of environmental temperature on lithium serum levels. Bipolar Disord. 2007;9(6):603–8.

Wright A, Sittig DF. Automated development of order sets and corollary orders by data mining in an ambulatory computerized physician order entry system. AMIA Annu Symp Proc AMIA Symp. 2006;2006:819–23.

Zarin DA, Young JL, West JC. Challenges to evidence-based medicine: a comparison of patients and treatments in randomized controlled trials with patients and treatments in a practice research network. Soc Psychiatry Psychiatr Epidemiol. 2005;40(1):27–35.

Zivanovic O. Lithium: a classic drug—Frequently discussed, but, sadly, seldom prescribed! Aust N Z J Psychiatry. 2017;1:4867417695889.