Longer Duration of Untreated Psychosis is associated with poorer outcomes for patients with Delusional Infestation
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ABSTRACT

We examined the association between the duration of untreated psychosis (DUP) and outcome for patients with delusional infestation (DI). This multi-centre international study included consecutive 211 patients. Illness severity was evaluated at first presentation and outcome was measured with the Clinical Global Impression scale (CGI) at base line and follow-up. A regression analysis showed a clear clinical and statistically significant association between shorter duration of untreated psychosis and better outcome at follow up. Patients with a DUP of less than 1 year showed a CGI-S change from 5.37 to 2.07; those with a DUP of 1-5 years a change from 5.48 to 2.59, and those with a DUP of above 5 years a change from 5.59 to 3.37. This difference of 1.1 CGI points between the groups resembles a clinically relevant difference in patient outcome Our results suggest that longer duration of untreated psychosis in patients with DI is associated with significantly less favourable clinical outcomes.

Key words: delusional infestation, duration of untreated psychosis, early intervention, outcome, clinical relevance, liaison psychiatry
INTRODUCTION

Delusional infestation (DI) is a psychiatric disorder in which patients have the delusional belief that they are infested with parasites or other living creatures (worms, fungi etc.), or inanimate pathogens such as fibres, threads or particles (1). DI is a psychotic disorder distinctly different from schizophrenia, schizoaffective and brief psychotic disorders. It is categorized as a delusional disorder, somatic type (297.1 in DSM-5; F22 in ICD-10). DI patients fail to fulfill criteria for schizophrenia, as they do not normally show disorganised speech, disorganised or catatonic behaviour, and negative symptoms such as blunting of affect, poverty of speech and thought, reduced social drive, loss of motivation, lack of social interest, and inattention to social or cognitive input. Hallucinations in DI, if present, are not prominent and related to the delusional theme of the infestation. Apart from the impact of the delusion(s) or its ramifications, functioning is not markedly impaired.

The prevalence of DI is estimated between 5.58 and 83.23 cases per 1 million inhabitants (2,3). Because patients with DI do not believe that they have a psychiatric illness, they usually seek referral to dermatologists or other specialists. Patients may also contact pest control businesses in order to detect and eradicate the perceived pathogen (1). Patients’ search for an identifiable infestation leads them to visit multiple physicians (1). As a result long duration of untreated psychosis (DUP) may be a common problem in patients with DI.

DUP is defined as the time that passes from manifestation of the first psychotic symptom to initiation of adequate evidence-based antipsychotic drug treatment (4,5). Defining the precise onset of psychosis can sometimes remain approximate as it may depend on patient recall. Nevertheless, generally, the measurement of DUP has proven to have good to excellent inter-rater reliability, as pointed out by Rubio and Correll. In their review they found that although heterogeneous definitions of DUP can complicate comparing DUP across studies, measurements of DUP are reliable within the same study (6). Furthermore, DI patients usually have a good memory for the time of perceived symptom onset because of the impairment of quality of life the symptoms normally cause. There is evidence of an association between long DUP and adverse clinical outcome in patients with schizophrenia (7-10). To
date, DUP and clinical outcome for patients with DI has not been systematically investigated. Available literature data on disease duration in DI are summarized in Table I. In publications disease/symptoms duration is equivalent to DUP, as patients are introduced to treatment with antipsychotics typically not earlier than from the moment of correct diagnosis.

Data on DI symptoms duration report disease durations varying from days to decades (11-23). For details see Table I. It has been noted that the duration distribution followed an exponential function with 52% of all cases showing a DUP of one year or less (24,25). Focusing exclusively on publications from the last 30 years, mean psychosis duration before receiving adequate treatment in DI varies from 0.97 to 5.6 years (see Table I) (1,18-20,22, 26-30). Only two studies provide mean duration values in a “M ± SD” format (23,30). The variance of median psychosis duration is less prominent. It ranges from 1 to 3.7 years (20,21,29,30,31), but the most frequently provided median duration is 1.0 year (22,24,26,27).

Some of the studies mentioned above include data on DI outcome. In these studies adherence to treatment was assumed but not verified, and DI patients are unlikely to have had a high adherence rate (32,33).

There are some suggestions that a short preclinical course in DI may indicate better outcome (24). However, the only study showing that shorter duration of DI was related to improved outcome had a rather small sample size (23 subjects) (29). Boggild et al. (30) showed that patients with full recovery or symptoms described as mild had a shorter DUP than patients with an incomplete or absent remission.

The aim of the current international multicentre study was to verify mean duration of untreated psychosis in patients with delusional infestation. Our hypothesis is that longer duration of untreated psychosis is associated with poorer outcome as assessed by change in clinical presentation during treatment. We arrive at this hypothesis based on similar associations in other mental illnesses between longer DUP and poorer outcome.

**METHODS**
Our consecutive cohort study comprises 211 cases of DI seen in multidisciplinary outpatient clinics in England (London 89 patients, Liverpool 32), Italy (Bruneck, 26) and Russia (Moscow, 64) over a period of 10 years (2006–2015). The included clinics specialized in the treatment of DI. These clinics receive referrals from a wide range of sources. Most cover large geographical areas. All clinics consist of a psychiatrist and a physician, either a dermatologist or a specialist in tropical medicine. In setting up our study, we established that all patients underwent a similar clinical approach. This includes the staffing of the clinics, follow up intervals according to clinical need, and the use of second generation antipsychotic medication as first line treatment. We excluded centres which were not able to demonstrate clinical uniformity. Standard clinical psychiatric and dermatological assessments were carried out for all patients to establish a diagnosis of DI. There were no patients identified with a genuine infestation. Once the diagnosis of DI was established, all patients were offered antipsychotics as treatment. In addition, they were offered appropriate treatment for any secondary skin condition or other secondary illness that may have triggered their DI. All patients were followed up rigorously as much as possible. Data on age and sex at presentation, duration of untreated psychosis (DUP), and disease severity (CGI severity score at baseline and CGI severity score at last follow-up) (34) were obtained. DUP was defined as the time from manifestation of the first psychotic symptom to initiation of evidence-based adequate antipsychotic treatment (35).

The severity of the DI was measured with the Clinical Global Impression Severity Subscale (CGI-S), a well-validated physician assessed scoring tool used frequently in psychiatry which grades symptom severity from 1 to 7, 1 representing “normal, not at all ill”; 2 representing “borderline mentally ill”; 3 representing “mildly ill”; 4 representing “moderately ill”; 5 representing “markedly ill”; 6 representing “severely ill”; and 7 representing “among the most extremely ill patients” (34). CGI-S was assessed at baseline and last follow-up.

**Statistical analysis**

Simple demographic frequencies are presented to identify the general characteristics of the population. To understand possible selection bias, we compared completers and dropouts with regards to those characteristics. We then investigated change between CGI-S at baseline and follow up by means of a
paired sampled t-test. In addition, an analysis of variance (ANOVA) was performed. We may expect DUP to be positively skewed (24). For this reason, we first categorized DUP in three groups in line with the literature (35, 36), those below and above 1 year and the outliers with a DUP of above 5 years. In accordance with the literature these groups have clinical relevance: we may expect treatment compliance to be better in the first as opposed to the last group (36). The ANOVA was performed to investigate group differences of CGI-change between these three clinically relevant groups. Following this, we performed a linear regression to investigate if one of the background characteristics other than duration of untreated psychosis were related to the treatment response as measured by change in the CGI. We first performed a Pearson and, where necessary, Spearman correlation to understand the direct association between the predictors age, sex, country and DUP and CGI change. We then included these variables in a multivariable (37. 38) regression, using the relevance criterion of Braun & Oswald, 2011 (39). A stepwise, forward entry, backward deselection linear regression procedure was used to identify variables also associated with CGI-S change. To adjust for DUP skewedness, we repeated the analysis, using log transformed DUP as predictor, in order to improve the understanding of the effect of the variable “DUP” on outcome (38). This careful procedure provides an impression of the contribution of possible confounders in the final model.

RESULTS
The consecutive sample consisted of 211 patients (Table II). In this sample the mean age was 58.8 years, with a minimum of 18 and a maximum of 95 years. 65% (n=138) of patients were female. The mean duration of untreated psychosis (DUP) was 3.4 years (SD 4.2), with a minimum of 2.5 months and a maximum of 29 years. The median was 2.0, with a positive skew value of 2.8, implying a number of outliers at the longest number of years of DUP. 37 patients did not engage with treatment or were lost to follow up, 1 patient died from an unrelated illness. The majority of patients (n=173, 82.0%) accepted treatment and confirmed relatively higher adherence to treatment matching previously published data from multidisciplinary psycho-dermatology clinics (32). Age, sex distribution, CGI-S at baseline and DUP in the patients refusing treatment did not differ from those accepting treatment. These 173 were included in the comparative analyses. 74 were from London, 57
from Moscow 24 from Bruneck and 18 from Liverpool. Treatment dropout differed significantly across countries, with the largest dropout seen in England, and the lowest in Italy. Figure 1 shows the distribution of DUP across all included patients.

Correlations and Analysis of variance

Table III shows the main result with the numbers of patients per DUP category and parametric analysis of variance comparing CGI-S scores at baseline and follow up. This shows a clear and positive association between longer DUP and poorer outcome. The table illustrates that in patients with a DUP above one year, and more importantly in the outliers, mean scores and mean ranks are substantially and significantly higher. When we looked at the DUP changes in those three groups with an ANOVA, patients with a DUP of less than 1 year showed a CGI change of 3.30 on the seven point CGI-S scale (CGI-S change from 5.37 to 2.07). Those with a DUP of 1-5 years showed a CGI-S change of 2.89 points (5.48 to 2.59), while those with a DUP of above 5 years showed a change of 2.22 CGI-S points (5.59 to 3.37). There was no statistical difference between the three groups with regards to the CGI-S scores at baseline (p=0.610). However, the difference between the groups was significant at follow-up (p=0.010), as was the significance of the difference of the change in CGI-S scores (p=0.037).

Our analysis of CGI-S score changes from baseline to last follow-up revealed a marked improvement in symptoms (mean change from 5.46 to 2.64; table II). Whereas before treatment most patients were on average ‘markedly ill’, with treatment the median score was ‘borderline ill’. A paired sample t-test between CGI-S scores at baseline and at follow-up showed a statistically significant difference (mean difference=- 2.87; t-value=22.37, p<0.001; table II). Importantly, the median reduction was 3 points in CGI-S severity, which is highly clinically relevant.

We made these detailed results visually more accessible in Figure 2, showing the relationship between DUP (in years) and CGI-S change. Figure 2 shows a downward slope of DUP the larger the CGI-change. This means that in general there was more improvement of CGI with shorter time of DUP. Furthermore, we categorized the patients into three clinically important groups according to the CGI severity at follow up: (1) patients not at all ill (n=62; 35%, CGI-S: 1), (2) patients borderline or mildly
ill (n=70; 40%, CGI-S: 2-3) and (3) patients moderately ill or above (n=41, 25%, CGI-S: 4-7).

Between these three groups, the mean DUP differed significantly, with a mean DUP of 2.18 in the ‘not at all ill’ group (group 1), 3.9 in the ‘borderline or mildly ill’ group (group 2) and of 4.22 in the ‘moderately ill’ or above group (group 3) (f-value=3.83, p=0.024).

When examining the change in severity from baseline, we identified three clinically important groups: (1) those much improved with a CGI-S score difference of more than four points, (2) those somewhat improved (CGI-S score difference of 1-3 points), (3) those with unchanged CGI-S scores or worsened CGI-S scores (change of 0 to -1 CGI-S points). Between these groups, the mean DUP differed significantly, with a mean DUP of 2.54 in the much improved, of 3.66 in the somewhat improved, and of 5.10 in the unchanged or worsened group (f-value=3.24, p=0.041).
Correlational and regression analysis in order to investigate confounders

A regression analysis was performed in a number of steps (Table IV and V). To identify relevant predictors, we first performed a Pearson and Spearman correlation (Table IV). Age was left out of the regression, as it showed a correlation of 0.011 to CGI-S change. In the regression analysis (Table V), DUP proved to be a significant predictor after correction for country as a confounder. The final model showed that gender was not related to changes in CGI-S scores, whilst country (Beta=0.705, 95% CI=0.514 to 0.896; p<0.001) and duration of untreated psychosis (Beta=0.068, 95 CI=0.014 to 0.122; p=0.014) showed an association. Using log transformed DUP as a predictor (38) showed that country (Beta=0.356, 95 CI=0.405 to 0.927; p<0.001) and duration of untreated psychosis (Beta=0.208, 95 CI=0.108 to 0.546; p=0.004) had comparable associations. The adjusted R-square (as indicator of model effect) was 0.136 (ES\textsubscript{Dup}=0.157), and 0.146 (ES\textsubscript{Log Dup}=0.172) in these final models, which is reasonable, given the number of predictors and the sample size. Stratification of the analyses by country improved the explained variance to 0.186 and 0.312. Co-linearity diagnostics performed beforehand showed no underlying associations of variables. DUP was not confounded by country (table II).

These analyses show that in the final model a clear prediction of change in CGI-S by DUP is observed. It confirms that there is an association between longer DUP and poorer outcome as measured by change in CGI-S. For results of further analyses please contact the authors.

DISCUSSION

Our results indicate that there is a clinically relevant association between shorter DUP and increased improvement of CGI-S scores. Longer DUP was associated with significantly less symptomatic improvement (as measured by CGI-S scores). The mean duration of untreated psychosis (DUP) obtained in our study was 3.4 years and falls into the range of mean values published elsewhere (1,18-20,22,26-28), including studies with large samples sizes (1,24,28,29). Only small studies with small sample sizes have so far presented DUP of below three or above four years with the exception of Foster et al. (n=147, DUP: 2.6 years) (22). In general, most previously published mean values are within the standard deviation of our mean value. Thus, the population in our study could be considered
as representative of the population of DI patients described elsewhere.

The range of disease duration shows similarities with previous studies, i.e. a minimum of 2.5 months and a maximum of 29 years in our study. This wide range of DUP is comparable to previously published data (18,19,22,23).

The negative effects of a long DUP have been explored in numerous studies for a variety of psychiatric diagnoses (6,10,38,39-44). They are heterogeneous but all concluded that DUP is an important modifiable indicator of prognosis.

However, there are no data on DUP and its association with outcome in delusional disorders. There is a single comparative study that suggests that DUP in patients with delusional disorders does not differ significantly from DUP in patients with schizophrenia (46). Our results are consistent with the only previous small study on DUP in DI (30), showing that shorter duration of DI is related to improved outcome and prognosis. However, probably as a result of small sample size, the authors of that study could distinguish only two relevant follow-up groups: “full recovery/mild residual symptoms” versus “incomplete /absent remission” that differed significantly (1.2 (SD 0.8) vs. 3.0 (SD 1.5) years). In contrast, our results comprise more distinct strata (“much improved”, “somewhat improved”, “unchanged”, “worse”), the use of a standardized instrument for severity measurement (CGI-S) and a much bigger study sample (173 vs 23 subjects). In addition, our study is a truly consecutive, multicenter study which has followed up patients systematically from specialist multidisciplinary clinics.

**Clinical relevance**

Our findings have clear clinical relevance. The potential of DUP being modifiable raises the possibility of improving clinical outcomes by shortening DUP. In designing interventions to shorten DUP, it is important to identify factors contributing to DUP. Factors previously associated with a longer DUP include stigma-related concerns (47,48), an insidious mode of onset (49,50), and a diagnosis of non-affective psychosis compared with affective psychosis (51,52,53). In contrast, DUP shortening is associated with development of early interventions that reduce treatment delay and promote recovery as it has been shown in schizophrenia studies. Early Intervention is followed by an
improvement in the prompt treatment of people with first episode psychosis (54). It remains unclear at this point how applicable the principles of early intervention are for patients with DI. However, in keeping with general early intervention approaches of destigmatization and insight improvement in psychosis, educational and contact interventions may be potentially beneficial. Any early intervention programs for DI should be provided in partnership between mental health professionals and other physicians (“joint care”), as there is evidence for Joint clinics in DI (1). There is an opportunity not only for combined assessment and treatment, but also for cross-education between representatives of different medical specialties to improve care and thus DUP. This can be provided in combined psychiatric and dermatological or psychiatric and tropical medicine clinics with specialists able to address the question how to persuade the DI patient with little or no insight to shorten the period of untreated psychosis by trying evidence-based antipsychotic treatment.

Limitations

This study investigated patients from specialist settings. These settings are probably the optimal treatment venues currently available for patients with DI, as they provide combined dermatology or tropical medicine and psychiatric services (54). It is not clear whether the same association between DUP and outcome applies to patients being seen only in primary care or for patients with DI managed in general dermatology clinics without specialist psycho-dermatology expertise. However, given that even in specialist psycho-dermatology clinics, there is an association between DUP and poorer clinical outcomes, and given that there is evidence (albeit non-randomized) that care of patients with DI is optimized in specialist psycho-dermatology clinics, the association between DUP and poorer clinical outcome may be even worse in non-specialized clinics. Even in our specialist settings 18% of patients did not agree to try any medication, which is in keeping with earlier publications (31). Clearly this is not an inconsequential number. It highlights the difficulties with engagement of this challenging patient group.

Even though we had the biggest sample size of any such study to date and the model required 150 patients to yield statistically relevant results, bigger sample sizes may improve the results in future research. We therefore need to emphasize the clinical rather than the statistical relevance of the
association we found. Furthermore, general demographic factors as education, having a job, or being married may increase the power of the findings, as may possible drug abuse, which was examined by our group in a separate study that included patients of this sample (55).

We have not factored in the length of follow-up in our study and different length of follow-up could potentially influence outcome. We have found a statistical association, which is of course no proof of causality.

We did find a variation between countries. However, this variation was expected because of the differences in clinic settings. The Bruneck sample was responsible for much of the variance. It was by far the smallest and the only rural sample. It showed better follow-up rates, comparable baseline CGI and lower follow-up CGI than the urban samples. This effect has been well documented in the literature where treatment compliance and effect is often better in rural versus urban settings (56).

**Suggestions for future research**

Research in DI is difficult as few randomized controlled trials exist for a variety of practical and other issues (2). Possible factors that contribute to longer DUP in DI should be studied. It will be essential to identify which barriers the groups with the shortest DUP and longest DUP had to overcome before starting treatment. This would require a qualitative research approach examining a variety of treatment approaches to identify the most successful approach in these patients. This may also include looking at characteristics of patients who refuse treatment. An interesting question for further studies may be whether early intervention (EI) models proposed for other psychoses are an adequate option for DI patients. A recent study showed that the concept of EI may be relevant in the age range of over 35-year-olds (57), more akin to the DI population. Finally, assessing the length of DUP and clinical outcomes for patients with primary compared to secondary DI would be an important next step.

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*Declaration of Interest*

The authors report no potential conflicts of interest.
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\textsuperscript{a} Mean and median disease duration was calculated because primary data was published for every case.
Table II  Baseline data and description of the population

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<td>Moscow (Russia)</td>
<td>56 (87.5%)</td>
<td>8 (12.5%)</td>
<td></td>
</tr>
<tr>
<td>London and Liverpool (UK)</td>
<td>92 (76.0%)</td>
<td>29 (24.0%)</td>
<td></td>
</tr>
<tr>
<td>DUP by country : mean (SD)</td>
<td></td>
<td></td>
<td>0.967</td>
</tr>
<tr>
<td>Bruneck (Italy)</td>
<td>3.00 (5.91)</td>
<td>0.21</td>
<td></td>
</tr>
<tr>
<td>Moscow (Russia)</td>
<td>3.51 (4.45)</td>
<td>3.58 (4.91)</td>
<td></td>
</tr>
<tr>
<td>London and Liverpool (UK)</td>
<td>3.42 (3.87)</td>
<td>3.49 (2.91)</td>
<td></td>
</tr>
<tr>
<td>CGI – S baseline</td>
<td>5.46 (0.92)</td>
<td>5.29 (1.01)</td>
<td>0.262</td>
</tr>
<tr>
<td>CGI – S follow up</td>
<td>2.64 (1.77)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CGI – Change</td>
<td>-2.87 (2.02)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P (paired t test baseline – follow up)</td>
<td>&lt; 0.001</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Significant difference: chi-square=7/75, df=2, p=0.021

**Notes**
This table shows completers and drop outs are largely the same. Also no important difference is observed in DUP over countries either in the completers (n=173, df=2, f=0.2, p=0.888) or in the non-completers (presented above, n=38, df=2, p=0.635). Italy is difficult to compare to the other locations due to the low numbers.
An analysis of variance of change in CGI showed a significant (n=173, df=2; F=3.35; p=0.037) difference in group means between the three DUP categories <1 year (μ=3.30 sd=1.66), 1 to 5 years (μ=2.89 sd=1.64) years and above 5 years (μ=2.20 sd=1.80) over change in CGI severity.
Table III  
Numbers of patients per DUP category and analysis of variance (baseline, follow up and change of CGI-S)  a

<table>
<thead>
<tr>
<th></th>
<th>CGI–S baseline</th>
<th>CGI–S follow up b</th>
<th>CGI–S change b</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n=</td>
<td>mean score</td>
<td>mean score</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(ANOVA)</td>
<td>(ANOVA)</td>
</tr>
<tr>
<td>DUP &lt; 1 year</td>
<td>41</td>
<td>5.37</td>
<td>2.07</td>
</tr>
<tr>
<td>DUP &gt; 1 year and &lt; 5 years</td>
<td>105</td>
<td>5.48</td>
<td>2.59</td>
</tr>
<tr>
<td>DUP &gt; 5 years</td>
<td>27</td>
<td>5.59</td>
<td>3.37</td>
</tr>
<tr>
<td>P=</td>
<td>173</td>
<td>0.610</td>
<td><strong>0.010</strong></td>
</tr>
<tr>
<td>CGI score all completers</td>
<td>173</td>
<td>5.46</td>
<td>2.64</td>
</tr>
</tbody>
</table>

Notes.

a We have applied a number of different statistical models to test whether there is an association between DUP and outcome. All models point in the same direction confirming that such an association exists. The regression may be seen as a confirmation of the non-parametric correlation and the parametric analysis of variance, which are the most important findings.

b We have used changes in CGI-S scores rather than CGI-Improvement (CGI-I) scores. This has the advantage of improving the statistical accuracy of the data but changes in CGI-S scores do not directly translate into CGI-I scores.
Table IV  Correlation of CGI-S change with age, gender, country and DUP (n=173 treatment completers)

<table>
<thead>
<tr>
<th>Predictors</th>
<th>Pearson correlation</th>
<th>P</th>
<th>Spearman correlation</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>0.011</td>
<td>0.899</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sex</td>
<td>0.135</td>
<td>0.077</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Country</td>
<td>0.316</td>
<td>.001</td>
<td></td>
<td></td>
</tr>
<tr>
<td>DUP</td>
<td>0.129</td>
<td>0.090</td>
<td>0.111</td>
<td>0.146</td>
</tr>
<tr>
<td>Models</td>
<td>Predictors</td>
<td>β</td>
<td>SE β</td>
<td>Standardized β</td>
</tr>
<tr>
<td>-------</td>
<td>------------</td>
<td>---------------</td>
<td>--------</td>
<td>----------------</td>
</tr>
<tr>
<td></td>
<td>Gender</td>
<td>0.466</td>
<td>0.267</td>
<td>0.132</td>
</tr>
<tr>
<td></td>
<td>DUP</td>
<td>0.069</td>
<td>0.029</td>
<td>0.178</td>
</tr>
<tr>
<td></td>
<td>Country</td>
<td>0.705</td>
<td>0.162</td>
<td>0.316</td>
</tr>
<tr>
<td>Model 1</td>
<td>Gender</td>
<td>0.300</td>
<td>0.315</td>
<td>0.071</td>
</tr>
<tr>
<td></td>
<td>Country</td>
<td>0.672</td>
<td>0.165</td>
<td>0.301</td>
</tr>
<tr>
<td>Model 2</td>
<td>Gender</td>
<td>0.283</td>
<td>0.314</td>
<td>0.067</td>
</tr>
<tr>
<td></td>
<td>Country</td>
<td>0.677</td>
<td>0.165</td>
<td>0.299</td>
</tr>
<tr>
<td></td>
<td>DUP</td>
<td>0.054</td>
<td>0.165</td>
<td>0.119</td>
</tr>
<tr>
<td>Model final</td>
<td>Country</td>
<td>0.705</td>
<td>0.160</td>
<td>0.339</td>
</tr>
<tr>
<td></td>
<td>DUP</td>
<td>0.068</td>
<td>0.188</td>
<td>0.162</td>
</tr>
</tbody>
</table>

* We used the uncategorized (skewed) DUP as predictor. Age was left out of this analysis, as it showed no association to CGI change in the correlational analysis. The other variables showed an association with a significance level below 0.2 and were included (39). This was repeated with log transformed DUP as predictor, to investigate the impact of the extreme skewedness of the variable on CGI change as outcome.

a In this procedure the predictors are first analysed separately. Then each variable is added to the model to identify the contribution of each variable to the final model, corrected for the other variables. Finally, a backward deselection was performed, deleting each item with a p < 0.2 from the full model.

b The beta – coefficient represents the effect of change in the predictor (age, gender, country or DUP) on outcome (CGI-S change). It is a measure of effect size, where 0.1 is small, 0.3 is medium and above 0.5 is large.

c Interpretation adjusted R square: R is the correlation between the predicted values and the observed values of Y. R square is the square of this coefficient and indicates the percentage of variation explained by the regression line out of the total variation. This value tends to increase as you include additional predictors in the model. Thus, one can artificially get higher R square results by increasing the number of Xs in the model. To rectify this effect, adjusted R square is used. When comparing models one should rely on adjusted R-square. This means that if R-square (adjusted) is 0.136, your model accounts for 13.60% of the total variability. The corresponding effect size Cohen’s f (R²/(1 - R²)) was 0.157, which is small. When using the logarithm of DUP, the R-square (adjusted) was 0.146, The corresponding effect size Cohen’s f (R²/(1 - R²)) was 0.172, which is reasonable.
Figure 1: Distribution of DUP
Figure 2:
Relationship between DUP (in months) and CGI-S score change*

* In the figure we only present the mean DUP per change category.