Trajectories of Treatment Response in Hallucinations

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ABSTRACT

Objective
Hallucinations are highly prevalent in schizophrenia and related disorders. Antipsychotics are generally effective in treating hallucinations, but major inter-individual differences in treatment response exist. Previous studies have identified heterogeneity of over-all antipsychotic response patterns in schizophrenia. The aim of this study was to explore the heterogeneity in the response of hallucinations to antipsychotic drug treatment in a representative sample of patients acutely admitted for psychosis.

Methods
226 adult patients with symptoms of active psychosis were included in a randomized pragmatic trial of second-generation antipsychotics and followed for 27 weeks. Latent-mixture and latent growth curve models were conducted to analyze heterogeneity of treatment response for hallucinations to second-generation antipsychotics.

Results
We found five different trajectories of treatment response for patients with hallucinations at baseline. These included two groups of “dramatic responders” who had rapid reduction followed by extinction of hallucinations during the first four weeks of treatment, then groups of “gradual responders”, “temporal responders” and “non-responders”. Most responders, 80% of those with hallucinations at baseline, were dramatic responders. Patients who showed no response in the early weeks remained non-responders also after longer follow-up.

Conclusions
The study suggests the existence of differential response patterns of hallucinations to antipsychotic treatment, and that a significant subgroup are dramatic responders. Hallucinations generally respond quickly to antipsychotic treatment. With no improvement in the very first weeks an early change of treatment should be considered.

Keywords
Hallucinations, antipsychotics, treatment response, trajectories, psychosis, heterogeneity, schizophrenia
Introduction

Response to antipsychotic drugs is typically reported dichotomously as rates, and considerable variations are found in recent meta-analyses depending on the specific response criteria used [1,2]. However, this responder/ non-responder dichotomy does not capture the range of responses that can be observed in clinical care. A different approach has been to investigate the heterogeneity of antipsychotic response patterns, where different response trajectories have indeed been identified across studies [3-8]. Several issues related to sample characteristics and methodological considerations applied. First, the majority of trajectory studies include patients with chronic schizophrenia which are unlikely to be generalizable to first episode and early-phase patients (8). Secondly, most studies include selected samples, use of experimental designs at odds with usual clinical practice and short study durations which may further limit generalizability [9]. Finally, most if not all previous studies use joint total scores of several symptom clusters or sum scores of single symptoms clusters as outcome measures [10,11]. By focusing on sum scores there is a risk to overlook the heterogeneity in treatment response in any particular symptom, as this might be cancelled out by opposing changes in other symptoms. A new approach would be to focus on a single symptom (see [12] for a discussion of single symptom approach in psychiatry research). Although sharing the same reality breach, psychotic disorders are heterogeneous symptomatically as demonstrated in the various diagnostic manuals [13,14].

Hallucinations are prevalent in schizophrenia and related disorders [15]. As much as 80% of patients with schizophrenia experience hallucinations where auditory hallucinations are most frequent [15]. Hallucinations may cause substantial suffering and are sometimes associated with suicidality [16-19], violence [20,21] and homicide [22]. Hallucinations are therefore symptoms of both major clinical importance and with distinguishing properties.

We have previously demonstrated that differential effectiveness may exist among second generation antipsychotics for hallucinations using a pragmatic RCT with a 24-months follow-up period [23]. Corresponding findings were also reported in a larger study [24]. An important new research question relates to whether different response trajectories of hallucinations exist independently of the particular antipsychotic drugs used.

The aims of the study were to investigate the heterogeneity of hallucinatory response to antipsychotic drug treatment in a representative sample of patients acutely admitted for psychosis.

Material and Methods

Study design

The present study uses data from the Bergen Psychosis Project (BPP), a 24-month prospective, randomized, pragmatic antipsychotic drug trial. The design and methods have been described in more detail elsewhere [25]. The study was conducted between 2004 and 2009 at the Division of Psychiatry, at Haukeland University Hospital in Bergen, Norway. The hospital serves a catchment population of about 400,000. The project was approved by the Regional Committee for Medical Research Ethics n Western Norway (REK Vest) and the Norwegian Social Science Data Services (NSD). The Regional Committee for Medical Research Ethics allowed eligible patients to be included before informed consent was provided, which facilitates a clinically relevant representation in the study. The BPP was publicly funded and did not receive any financial or other support from the pharmaceutical industry. The present study focuses on the first six months of follow-up.

Sample

The sample was consecutively recruited from the psychiatric emergency ward. Adult patients were eligible for the study if they were admitted for symptoms of active psychosis as determined by a score of ≥ 4 on one or more of the items Delusions, Hallucinations, Grandiosity, Suspiciousness/persecution or Unusual thought content in the Positive and Negative Syndrome Scale (PANSS) [26] and were candidates for oral antipsychotic drug therapy. All included patients met the diagnostic criteria of the International Classification of Disease ICD-10 [13] for schizophrenia, schizoaffective disorder, acute and transient psychotic disorder, delusional disorder, drug-induced psychosis, major depressive disorder with psychotic features and bipolar disorder except manic psychosis. Patients with drug-induced psychosis were only included when the condition did not resolve within a few days and when antipsychotic drug therapy was indicated. The diagnoses were determined by the psychiatrists or specialists in clinical psychology in the hospital. Exclusion criteria were inability to use oral antipsychotics,
inability to cooperate during the investigation because of manic psychosis or other behavioral or mental reasons related to the state of illness, not understanding Norwegian language, treatment with clozapine at admission or being candidate for electroconvulsive therapy treatment.

A total of 226 adult (> 18 years) patients were at inclusion randomized to one of the following antipsychotics: risperidone, olanzapine, quetiapine or ziprasidone. In accordance with a pragmatic design aiming to mimic everyday clinical decision making, the randomization was open to both the patient and the clinical staff, whereas the assessments were conducted by blinded research personnel. Furthermore, the patients were randomized to a random sequence of the investigational agents rather than a single drug. The patients were offered the first agent in the sequence but could choose the next agent in the sequence if the first drug could not be used because of contraindications or prior negative experience. The same procedure was followed if the next antipsychotic drug(s) in the sequence could not be used. The treating psychiatrist was free to choose the dosing, combination with other psychotropics and discontinuation and change to another antipsychotic agent in the presence of insufficient efficacy and/or side effects. In the rare case that a patient already used an antipsychotic agent in therapeutic dosage at admission, no wash-out was performed before starting the study drug. If a patient was randomized to the same agent already under use, the agent would simply be continued and only dosage at admission, no wash-out was performed used an antipsychotic agent in therapeutic effects. In the rare case that a patient already change to another antipsychotic agent in the other psychotropics and discontinuation and could not be used. The treating psychiatrist was the next antipsychotic drug(s) in the sequence if the first drug could not be used because of contraindications or prior negative experience. The same procedure was followed if the next antipsychotic drug(s) in the sequence could not be used. The treating psychiatrist was free to choose the dosing, combination with other psychotropics and discontinuation and change to another antipsychotic agent in the presence of insufficient efficacy and/or side effects. In the rare case that a patient already used an antipsychotic agent in therapeutic dosage at admission, no wash-out was performed before starting the study drug. If a patient was randomized to the same agent already under use, the agent would simply be continued and only dose adjustment made if indicated.

**Measurements**

The patients were assessed at baseline (T1), at discharge or latest at 9 weeks (mean 4.1 weeks) if not discharged earlier (T2) and at follow-up visits after 3 (T3) and 6 months (T4). The main reason for drop-out between T1 and T2 was discharge from the hospital before one week had passed since baseline, which was defined as the minimum interval before T2 assessments could be undertaken. There were no baseline differences in any clinical or demographic variables between those with baseline assessments only and those with follow up assessments, except for higher PANSS negative subscale score in those with baseline assessments only (independent samples t-test: p=0.02; mean difference 2.3; 95% confidence interval of the difference 0.4-4.2). The severity of hallucinations was evaluated by item P3 (Hallucinatory behavior) of the PANSS positive subscale, where a score of 3 or more indicate the presence of hallucinations. The Intraclass correlation coefficient (ICC) for the PANSS was calculated based on inter-rater assessments between EJ, HAJ and RAK, and showed high inter-rater reliability (0.92). Calgary Depression Scale for Schizophrenia (CDSS) was used to measure severity of depression [27]. CDUS/CAUS - the Clinical Drug and Alcohol Use Scales – were conducted for overview of the use of alcohol and illicit drugs [28]. The Clinical Global Impression-Severity of Illness scale (CGI-I) [29] and Global Assessment of Functioning-Split Version, Functions scale (GAF-F) were used to assess the general functioning level [30]. A brief neuropsychological screening instrument; the Repeatable Battery for the Assessment of Neuropsychological Status (RBANS), was administered at baseline [31,32]. Serum level measurements of antipsychotic medication were performed at all follow-up visits.

**Statistical data analysis**

SPSS 22 was used for descriptive statistics, frequency, cross tabulation with chi-square and group difference t-test [33]. Level and change at mean and individual level was analyzed with latent growth curve modeling (LGC) [34]. The outcome variable (PANSS item P3) was ordinal; however, with seven categories and therefore analyzed as a continuous variable. The Maximum Likelihood Estimator with robust standard errors (MLR) was used in order to give unbiased estimates when non-normality in data was experienced [35]. The Full Information Maximization Likelihood method uses all available data under the “Missing at Random” assumption [35,36], minimizes the negative effect of missing data, improves statistical power and generalizability of the results compared to models based on the ordinary listwise deletion method [37].

An LGC-model from T1 to T4 revealed estimation problems caused by the linearity restriction. A quadratic factor was added in order to describe a non-linear change over time. However, this model did not fit the data (CFI=0.58, TLI=0.37, RMSEA=0.18) and showed estimations problems as well. These results together with a visual inspection of the observed data showed further needs for improvements. In the final model one linear difference factor that described level and change from T1 to T2 after approximately four weeks (S: 0 – 4.1 weeks) and another latent factor
capturing the change from T2 to T4 (S2: 4.1-27.0 weeks) was estimated.

Then, Latent Class Growth Analysis (LCGA) was conducted to describe the heterogeneity in the data expressed as the maximal number of possible trajectories [38]. The evaluation of the number of classes was based on the entropy index and model fit indices Akaike Information Criterion (AIC), the Bayesian Information Criterion (BIC) and sample-adjusted BIC, with lower values indicating better model fit [35,39]. Statistically significant improvement of adding classes was tested by the Vuong-Lo-Mendell-Rubin Likelihood Ratio Test for k-1 versus k classes (LRT) [39]. The estimated class size and relative frequencies were used in this evaluation. A sample size of 25 classified subjects and 5% of the sample has been proposed as threshold values [40]. Mplus 7.4 was used for LGC analysis and LCGA [36].

Results

A total of 226 patients were included, of which 74 (32.7%) were females. The mean age at baseline was 34.1 (SD 13.5) and 44.2% had not used antipsychotic drugs before. The distribution of the study-drugs at baseline was 23.6% risperidone, 29.8% olanzapine, 21.8% quetiapine and 24.9% ziprasidone. At the last follow up the distribution of medications was similar.

The descriptive statistics and sample size for the outcome variable at baseline (T1), at first follow-up (mean 4.1 weeks after baseline) (T2) and at follow-up visits after 3 (T3) and 6 months (T4) are presented in Table 1. The use of antipsychotic drugs and concomitant psychotropic medications are displayed in the supplementary material, Supplementary Table 1.

- Level and change in hallucinations over time in the total group

Latent growth curve (LGC) analyses were used to investigate the level and change in the hallucination score for the total sample collectively. After exploring different models, one linear change factor that described level and change from T1 to T2 (S1: 0-4 weeks) and another factor capturing the change from T2 to T4 (S2: 4-27 weeks), were estimated. The mean baseline PANSS P3 score was estimated to be 3.51 (p<0.001) with individual variance=2.65 (SD 1.63), the mean change from T1 to T2 was -0.38 per week (p<0.001) and the latent change from T2 to T4 was -0.01 per week (p=0.06). The individual variance in change for the two slope factors were S1: SD=0.38 (p<0.01) and S2: SD=0.03 (p=0.51). Stronger reductions in the scores from T1 to T2 were seen in patients with higher baseline scores than in patients with lower baseline scores (r=-0.71, p<0.001). This model fitted data well ($\chi^2=2.63$, df=2, p=0.27, Comparative fit index (CFI)=0.99, Tucker-Lewis Index (TLI)=0.97, Root mean square error of approximation (RMSEA)=0.037, RMSEA CI=0.000-0.143, RMSEA close fit=0.45).

- Description of different trajectories

Latent class analyses (LCGA) revealed up to seven trajectory classes (Table 2). The LRT- and entropy results indicated three and seven classes as the best models. The smallest class in the three class model was larger in size than in the seven class model. However, the seven class model more precisely captured the individual variability in level and changes and was therefore chosen for presentations. The model is presented in Figure 1 and show trajectories with different baseline levels and changes of hallucinations: one high hallucination-level class that maintain the symptom level during the follow up (C1: 3% of the sample); one class with initially high hallucination scores with a temporary improvement and then worsening of the symptom level (C2: 4%); two large classes with early improvements (C3: 30%) and (C7: 24%); one low hallucination-level class with worsening of the symptom at 4 weeks and then improvement (C4: 3%); a low hallucination-level class that maintain the same level during the follow-up (C5: 29%) and finally a high hallucination-level class with slow, but progressive reduction of the hallucinations score (C6: 6% of the sample).

- Hallucinating subgroup

Three groups of patients with hallucinations at baseline had marked reductions of hallucination scores, of which two groups (80% of those with hallucinations at baseline) might be categorized as “dramatic responders”. This group had a rapid reduction followed by extinction of hallucinations during the first four weeks of treatment. One group – (10% of those with hallucinations) showed a more gradual but ultimately strong response and might be categorized as “gradual responders” One group (7% collectively of those with hallucinations) had temporary reduction of hallucinations and might be categorized as “temporal responders”. Non-responders (4% collectively of those with hallucinations) were patients who had no reduction of the hallucination score.
Table 1: Descriptive results for PANSS P3 – Hallucinations.

<table>
<thead>
<tr>
<th></th>
<th>N</th>
<th>Mean</th>
<th>SD</th>
<th>Skewness</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1</td>
<td>226</td>
<td>3.51</td>
<td>1.63</td>
<td>-0.43</td>
</tr>
<tr>
<td>T2</td>
<td>109</td>
<td>2.03</td>
<td>1.42</td>
<td>1.05</td>
</tr>
<tr>
<td>T3</td>
<td>52</td>
<td>1.73</td>
<td>1.29</td>
<td>1.51</td>
</tr>
<tr>
<td>T4</td>
<td>39</td>
<td>1.85</td>
<td>1.48</td>
<td>1.41</td>
</tr>
</tbody>
</table>

Table 2: Latent Class Growth Analysis (LCGA) of the PANSS hallucination indicator with fit measures for evaluation: Akaike Information Criterion (AIC), Bayes Information Criterion (BIC), Sample-adjusted BIC (SABIC) and Likelihood Ratio Test significance difference test (LRT).

<table>
<thead>
<tr>
<th>Classes</th>
<th>AIC</th>
<th>BIC</th>
<th>SABIC</th>
<th>LRT p-value</th>
<th>Entropy</th>
<th>Least class predicted n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1575</td>
<td>1599</td>
<td>1576</td>
<td>-</td>
<td>-</td>
<td>34 (15%)</td>
</tr>
<tr>
<td>2</td>
<td>1418</td>
<td>1455</td>
<td>1420</td>
<td>0.014</td>
<td>0.58</td>
<td>33 (15%)</td>
</tr>
<tr>
<td>3</td>
<td>1320</td>
<td>1372</td>
<td>1324</td>
<td>0.000</td>
<td>0.74</td>
<td>4 (2%)</td>
</tr>
<tr>
<td>4</td>
<td>1286</td>
<td>1351</td>
<td>1290</td>
<td>0.173</td>
<td>0.70</td>
<td>6 (3%)</td>
</tr>
<tr>
<td>5</td>
<td>1259</td>
<td>1338</td>
<td>1268</td>
<td>0.274</td>
<td>0.71</td>
<td>3 (1%)</td>
</tr>
<tr>
<td>6ᵇ</td>
<td>1226</td>
<td>1318</td>
<td>1233</td>
<td>0.012</td>
<td>0.73</td>
<td>6 (3%)</td>
</tr>
<tr>
<td>7ᶜ</td>
<td>1213</td>
<td>1319</td>
<td>1221</td>
<td>0.031</td>
<td>0.75</td>
<td>6 (3%)</td>
</tr>
<tr>
<td>8ᵈ</td>
<td>1148</td>
<td>1268</td>
<td>1157</td>
<td>0.270</td>
<td>0.71</td>
<td>4 (2%)</td>
</tr>
</tbody>
</table>

ᵇVuong-Lo-Mendell-Rubin Likelihood Ratio Test for k-1 versus k classes
ᶜUnstable solution. The model was reestimated with 10,000 starting values
ᵈConvergence problems: solution: Starts = 10000 400
ᵉConvergence problems: solution: Starts = 100000 500

Figure 1: Results from Latent Class Growth Analysis (LCGA) showing level and changes over time in hallucinatory behavior for seven classes/ trajectories (C1-C7). Frequencies for the most likely latent class membership are given.
Discussion

The main objective of the study was to disentangle different response trajectories of hallucinations as a prevalent and clinically relevant symptom. Seven different response patterns were identified, five of those included patients with hallucinations at baseline. More than three-quarters of patients with clinically significant hallucinations at baseline could be defined as dramatic responders, with extinction of hallucinations during the first four weeks of treatment. This shows that hallucinations tend to respond rapidly and well to antipsychotic medication in the vast majority of psychotic patients troubled by this symptom. In the shared decision process, this information is relevant to patients and caregivers.

Most studies that have investigated heterogeneity in treatment response, have focused on total psychopathology or one of the symptom sub-dimensions which complicates direct comparisons of our results to other studies. However, studies typically report four to five distinct response patterns. Marques and colleagues [5] found four response trajectories for positive symptoms during six weeks of follow-up. These are categorized as “dramatic responders” (patients that start off with a high symptom level and show dramatic decrease of symptoms), “responders” (patients that start off with average symptom scores and show robust response), “partial responders” (patients that show modest response) and finally “non-responders”. Pelayo-Teran and colleagues [11] found five trajectories for positive symptoms response during six weeks of follow-up. Comparable terminology is used among the studies for the different trajectories, including also a “slow partial responders” group in the latter study. Levine and colleagues [6] report response in total psychopathology during eight weeks of follow-up and found groups of “poor responders”, “rapid treatment responders” and three groups of “moderate responders”.

The proportions with non- or poor response are quite similar across studies, with 10% reported by Marques et al [5], 8.3% reported by Pelayo-Teran et al [11], 8% reported by Levine et al [6]. In our study non responders reached 7% after 6 month in the total sample and 11% in the hallucinating subgroup, however part of those patients were temporal responders, with rapid and significant response in the first 4 weeks, followed by gradual worsening of hallucinations. The group that could be characterized as “dramatic responders” was significantly larger in our study compared to the other studies [5,11]. Reasons for these differences may include a faster and more pronounced response of hallucinations compared to other symptoms of psychosis, such as for example delusions. Several other studies reports significantly faster response of hallucinations to antipsychotic treatment than that for delusions [41,42]. In this context, the delayed response of delusions to antipsychotic drug treatment would also influence the sum scores of positive symptoms in which delusions are embedded, causing the change of sum scores to appear at a slower rate.

Our results may be related to the pragmatic design of our study with a more clinically relevant sample compared to those in most RCTs. Indeed findings from the EUFEST study, a large pragmatic trial of antipsychotic effectiveness, found a substantially larger PANSS-score reduction compared to that reported in traditional randomized trials of antipsychotic efficacy [43], which among others include less diverse study populations because of numerous in- and exclusion criteria [44].

Results may also be related to our sample of patients in the acute phase of psychosis. Faster reduction of symptom would be expected in this patient group compared to more stable patients. Our study revealed a small but clinically important group of patients (3% of the total sample) who had sub-threshold scores for hallucinations at baseline, but then had an apparent worsening of symptoms, followed by a gradual decline and extinguishing of hallucinations. We can only speculate about the reasons for this particular response trajectory, but might include actual worsening of hallucinations during the acute phase or under-reporting of symptoms at baseline.

Results from our study might support the notion of different underlying mechanisms for the hallucinations. Findings in our study, where most of the patient are dramatic responders, might also support underlying pathophysiological dopaminergic mechanism, where high levels of dopamine in the limbic system play a major role, in most of the cases with hallucinations. However, non-responders might have a different underlying mechanism for the hallucinations [45] and other treatment options than D2-receptor blocking antipsychotics should be considered. Unfortunately, as of yet, not many other treatment options in the clinical care are
available as all of the antipsychotics primarily blocks D2 receptors (for discussion about personalized treatment of hallucinations see I. Sommer et al [45]).

Another important aspect of different response trajectories is their relation to clinical and non-clinical characteristics. Factors such as diagnosis, other symptoms, previous medication or different metabolism of the drugs, as well as gender, genetic variation, nutrition, specific diets or ethnicity might play a role in treatment response and thereby potentially serve as predictors of different response patterns [46-48]. Investigating such predictors was beyond the scope of this paper, however, but should be a focus in the future studies.

**Limitations**

The main advantages of the study are the consecutive recruitment of a clinically representative acute-phase sample and the focus on a single symptom of high prevalence and clinical significance, both of which are compliant with the recently suggested strategy for new drug developments in psychosis [49]. Some limitations should nevertheless be mentioned: High dropout rates are an important and well-known issue in antipsychotic drug trials [50]. The current study had a considerable attrition rate, but was comparable to other RCTs conducted on schizophrenia spectrum disorders [51]. Also, attrition did not seem to be associated with any baseline characteristics except a slightly higher PANSS negative subscale score in those with baseline assessments only. Choosing seven classes in LCGA model without reaching a proposed sample size of 25 classified subjects and five percent of the sample represent a limitation. We attempted to identify the highest number of highly homogeneous groups to represent the heterogeneity seen in every day clinical life, but because some of the classes were small, further statistical analyses of inter-class differences were not performed. These results need to be replicated in samples of larger size before any final conclusions can be drawn. Finally, hallucinations were investigated collectively as we did not have information beyond the hallucinations item of the PANSS scale. Theoretically, different modalities of hallucinations either might have individual response patterns, and future studies should include subtyping of hallucination modalities such as auditory, visual, tactile, and others.

**Conclusions**

This study contributes with new evidence concerning hallucinations as a separate target for antipsychotic drug treatment and a basis for stratified approaches. Hallucinations generally respond quickly to antipsychotic drug treatment and in the absence of response during the very first weeks of treatment, an early change of treatment should be considered. Future studies should aim at identifying predictors of the different treatment trajectories to better individualize treatment.

**Conflicts of Interest**

Iris Sommer is consultant to Gabather and has received research support from Janssen and Sunovion.

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**Author contributions**

All authors made substantive intellectual contributions to the study. IS planned the study, wrote the first draft of the manuscript and participated in the statistical analyses; RG undertook the statistical analyses, contributed in the interpretations of the data and helped draft the manuscript; EK contributed in the interpretations of the data and helped draft the manuscript; LR contributed in the interpretations of the data and helped draft the manuscript; KH contributed in the interpretations of the data and helped draft the manuscript; RAK collected data, contributed in analyses and interpretations of the data and helped draft the manuscript; EML contributed in analyses and interpretations of the data and helped draft the manuscript; HAJ designed the study, assisted in data collection, contributed in analyses and interpretations of the data and contributed to the drafting of the manuscript; EJ was the co-designer of the study, collected data, undertook the statistical analyses and helped draft the manuscript. All authors have read and given final approval of the latest version of the manuscript.
References


33. IBM Corp. IBM SPSS Statistics for Windows: Ver. 22.0. IBM Corp Armonk, NY (2013).


36. Muthèn LK, Muthèn BO. Mplus 7.3. 3463 Stoner Avenue, Los Angeles, California (2014).


42. Gunduz-Bruce H, McMenimam M, Robinson DG, et al. Duration of untreated psychosis and time to treatment response for delu-


