Aberrant salience relationship with first rank symptoms

Andrea Ballerini1*, Marta Tortorelli1, Paolo Marino1, Cristina Appignanesi1, Cinzia Baschirotto1, Luca Mallardo1, Tommaso Tofani1, Francesco Pietrini2, Giulio D'Anna1, Andrea Rossi1, Valdo Ricca1 and Marina Santella1

Abstract

Background: Aberrant salience is the incorrect assignment of salience, significance, or value to different innocuous stimuli that might precede the onset of psychotic symptoms. The present study aimed to perform a preliminary evaluation of potentially different correlations between the Aberrant Salience Inventory (ASI) score and dimensional or categorical diagnostic approaches.

Methods: 168 adult outpatients with a current psychiatric diagnosis were consecutively enrolled. Patients were evaluated using different psychometric scales. ASI was used to evaluate aberrant salience, and to evaluate the association between ASI scores and first rank symptoms (FRS), and/or with a psychiatric diagnosis. Principal dichotomic clusters of ASI were identified using the Chi-square automatic interaction detection (CHAID) method.

Results: Current (16.76 ± 6.02 vs 13.37 ± 5.76; p = 0.001), lifetime (15.74 ± 6.08 vs 13.16 ± 5.74; p = 0.005) and past (15.75 ± 6.01 vs 13.33 ± 5.80; p = 0.009) FRS were the main clusters dichotomizing ASI. The average ASI score did not significantly differ among patients with different diagnoses.

Conclusions: ASI could be used as a tool to identify psychopathological dimensions, rather than the categorical diagnoses, in the schizophrenic spectrum.

Keypoints

• Aberrant salience is a trans-nosographic trait of psychotic and pre-psychotic symptoms which is not commonly used in clinical practice.
• The use of the Aberrant Salience Inventory may constitute a valuable instrument for the early detection of psychosis.
• A dimensional approach evaluating aberrant salience may provide a different framework in the assessment of psychotic diagnoses.

Keywords: Aberrant salience, Psychotic symptoms, Dimensional approach, Schizophrenic spectrum

Introduction

The concept of salience was described by Kapur as “a process whereby objects and representations, through the process of association, come to be attention-grabbing and capture thought and behavior” [1]. Several studies suggested the main role of dopamine in this process [2–5]
as the mesolimbic dopamine system plays a central role in the attribution of salience [6], converting the representation of a neutral external stimulus into an attractive/aversive one [7]. On the other hand, the dopaminergic hyperactivation in the ventral tegmental area/substantia nigra and striatum is associated with the aberrant feelings of salience [8] and, if chronically maintained, can lead patients through psychotic-like experiences (PLEs) and delusional conclusions [9]. Patients’ PLEs are associated with a plethora of meaningful coincidences impinging on their current worldview, and every single event is perceived as pointing to a new reality. Eventually, delusions become a possible explanation to make sense of these experiences.

According to Kapur [5], patients with aberrant salience over-attribute the meanings of otherwise neutral stimuli, typical in the prodromal stages of psychoses. In fact, at this stage the onset of apprehension and anguish is common for a world that has become uncertain and full of new meanings: This “delusional atmosphere” has been described by Jaspers as “a change which envelops everything with a subtle, pervasive and strangely uncertain light” [10]. This hypothesis might link the aberrant signaling of motivational salience, through the dysregulated dopamine transmission [11], to psychotic symptoms, bridging the gap between neurobiology and phenomenology [1].

Questionnaires providing scores on the proneness to develop psychosis in the general population, including the Magical Ideation Scale [12], the Perceptual Aberration Scale [13], and the Referential Thinking Scale [14], are available. Starting from these scales, Cicero et al. created the Aberrant Salience Inventory (ASI) as a valid and reliable measure of aberrant salience in both the clinical and general population [9]. Even if the interest in this topic is growing, this test is rarely used in research on psychosis, or clinical practice. Furthermore, no study investigates whether aberrant salience is a state or a trait condition and whether aberrant salience assessment can identify proneness to psychosis.

Even if Schneider proposed using the first rank symptoms (FRS) as a reference for the psychiatric diagnostic process in 1959 [15], they are rarely used for clinical assessment; moreover, there is general uncertainty about their diagnostic accuracy [16]. In their review, Soares-Weiser et al. concluded that FRS could be used to help clinicians to assess patients’ clinical conditions, but they do not rely on them alone because FRS appear in the diagnostic criteria of schizophrenia and in other psychotic or non-psychotic mental disorders (FRS are included in the Diagnostic and Statistical Manual—DSM [17] and International Classification of Diseases—ICD [18] checklists). From a psychopathological point of view, psychotic symptoms could be identified in different psychiatric conditions, including affective and psychotic disorders. Using the categorical diagnostic approach, based only on DSM or ICD, the clinician risks misidentifying several psychiatric conditions, especially in their earliest stages, making it practically impossible to identify patients’ aberrant salience experience(s). This misidentification is likely to delay an early therapeutic intervention (psychotherapeutic, psychosocial, or pharmacological) in the prodromal stages of psychosis.

The present study aimed to perform a preliminary evaluation of the potential association between the ASI score with dimensional or categorical diagnostic approaches and, possibly, to identify psychotic proneness in a transnosographic real-world population.

### Methods

This is a cross-sectional, monocentric study, conducted by the psychiatric research team at the Adult Psychiatry Unit of Florence University Hospital, according to Good Clinical Practices (GCP), the Declaration of Helsinki (1964), and further revisions. The study protocol was approved by the Ethics Committee of the local institution. Before they were enrolled in the trial, all patients received an information leaflet, in Italian language, explaining exhaustively the protocol and its implications. Further explanations were given by the research team as needed. Before collecting personal data and evaluating clinical conditions, patients were asked to sign their written informed consent.

All outpatients evaluated at the outpatient facility of the Psychiatry Unit of Florence University Hospital, meeting the following inclusion criteria, were consecutively recruited between May and December 2019:

- Age between 18 and 80 years;
- Diagnosis of a psychiatric disorder (depressive disorders, obsessive–compulsive and related disorders, bipolar and related disorders, schizophrenia spectrum and other psychotic disorders, anxiety disorders, obsessive–compulsive and related disorders) according to DSM-V;
- Italian speakers;
- Elementary school degree or above scholarship.

The exclusion criteria were:

- Acute psychotic episode (according to clinical evaluation);
- Cognitive impairment due to psychiatric or medical conditions;
- Diagnosis of a neurological condition.
All diagnoses were assessed by expert psychiatrists (authors AB and VR). Only the main diagnosis was included in the database. The ASI questionnaire was answered by each patient together with a psychiatrist, to ensure the best questionnaire understanding and, thus, the reliability of data collection. One of the research team members was addressed to give any further explanation to patients and to collect socio-demographic and anamnestic data.

The following scales were assessed:

- Functional dimensions of the Manual for the Assessment and Documentation of Psychopathology (AMDP) [19];
- Aberrant Salience Inventory [9];
- Positive and Negative Syndrome Scale (PANSS) [20];
- Montgomery–Asberg Depression Rating Scale (MADRS) [21];
- Hamilton Anxiety Scale (HAMA) [22];
- Mania Rating Scale (MRS) [23];

The presence of Schneider’s FRS was assessed according to the following items in the Manual for the AMDP [24]:

- 33–46 for delusion,
- 47–52 for hallucinations,
- 50–53 for anomalous self-experience.

We have collected the presence of any FRS only, and analyzed it accordingly; AMDP items were not analyzed.

The ASI is a self-reported questionnaire including 29 “Yes”/“No” questions (“Yes”: one point; “No”: zero points), based on the phenomenological descriptions of the prodromal phase of psychosis [9]. According to Kapur’s conceptualization [25], this scale explores five aberrant salience domains:

- Sense sharpening (items 3, 9, 12, 18, and 22);
- Heightened cognition (items 4, 7, 13, 19, 23, and 25);
- Heightened emotionality (items 8, 14, 20, 24, 26, and 28);
- Increased significance (items 1, 5, 10, 15, 16, 21, and 26);
- Impending understanding (items: 2–6–11–17–29).

The Italian version was confirmed to be reliable to identify psychotic patients—who showed higher scores when compared with the general population [26, 27]. Only the total ASI scores were analyzed. An independent research student double-checked the final database, and any discrepancy was aligned with source paper records.

Data were anonymized to guarantee the confidentiality of patients, according to legal requirements, and entered into a computerized database (including personal data, diagnosis, presence of lifetime and current FRS, current pharmacological therapies, questionnaire scores). Only patients who had all ASI items collected were analyzed (per-protocol population).

A preliminary cluster analysis (Chi-square automatic interaction detection, CHAID) was conducted to investigate which of the collected variables dichotomized ASI at a statistically significant level; a step-by-step approach was used. All collected variables (clinical, demographical, psychopathological) were included to determine the first-order cluster discriminative variable. Variables identified as discriminative were excluded from the next steps. A similar analysis was performed on all psychopathological variables to confirm the reliability of the sample and the method. Correlations between total scores of assessed scales were checked.

Continuous variables were compared with Student’s t-test; categorical ones were compared with Chi-square.

Statistical significance was set at $p < 0.05$. All statistical analyses were conducted using the default setting of SPSS (Statistical Package for Social Sciences) for Windows 20.0.

**Results**

203 consecutive subjects were asked to enter the study, and 7 of them refused to give their consent. Overall, 196 patients were effectively involved in the study, but 28 participants did not fill one or more items of the ASI. 168 subjects completed the entire study protocol, and they were therefore included in our analyses. Socio-demographic and clinical data are summarized in Table 1.

The age of patients was between 18 and 77 years (mean: 44.38; SD: 15.09). A mood disorder diagnosis was the most common (58.3%). More than half of the patients (51.2%) had experienced at least one FRS during their life. Average HAM-A (12.27 ± 8.12) and MADRS (20.79 ± 12.99) scores highlighted a generalized depressive mood. Furthermore, PANSS scores (average PANSS total 58.08 ± 21.61; PANSS general 33.93 ± 12.53), together with the use of any antipsychotic in 92 subjects (54.2%), confirmed the high frequency of psychotic symptoms in this population (Table 2).

The complete comparisons of average ASI values are described in Table 3. Variables dichotomizing the total ASI score are shown in Table 4. Patients with FRS (current, lifetime, or past) had significantly higher ASI scores. When FRS were excluded from the cluster analysis, patients with MRS higher than 3 and PANSS positive symptoms above 7 had significantly higher ASI scores.
The cluster analyses also showed that depression diagnosis was the first-step discriminant for MADRS (24.98 ± 10.74 vs 18.80 ± 13.53; \( p = 0.004 \)) and HAMA (14.43 ± 7.51 vs 11.25 ± 8.23; \( p = 0.018 \)), the diagnosis of schizophrenia for PANSS negative symptoms (18.15 ± 9.78 vs 12.78 ± 7.06; \( p < 0.001 \)), while current FRS was the main discriminator for MRS (6.13 ± 4.60 vs 2.98 ± 3.41; \( p < 0.001 \)), PANSS positive symptoms (13.58 ± 6.15 vs 7.95 ± 1.90; \( p < 0.001 \)), PANSS general (38.40 ± 14.12 vs 31.75 ± 11.12; \( p = 0.001 \)) and PANSS total (69.53 ± 24.45 vs 52.51 ± 17.69; \( p = 0.000 \)). No discriminants were found for the other scales collected in this study.

The correlation between scales is summarized in Table 5. While correlation analyses consistently showed a good correlation between scales assessing the same or similar pathological dimensions, none of them showed a significant correlation with ASI.

### Table 1 Socio-demographic and clinical data

| Category                                | N  | %   |
|-----------------------------------------|----|-----|
| **Sociodemographic variables**          |    |     |
| Gender                                  |    |     |
| Female                                  | 95 | 56.5|
| Male                                    | 73 | 43.5|
| Educational level                       |    |     |
| Elementary school                       | 7  | 4.2 |
| Middle school                           | 39 | 23.2|
| High school                             | 93 | 55.3|
| University                              | 29 | 17.3|
| Marital status                          |    |     |
| Single                                  | 66 | 39.3|
| Married                                 | 51 | 30.4|
| Divorced                                | 42 | 25.0|
| Widow                                   | 9  | 5.4 |
| Occupation                              |    |     |
| Student                                 | 21 | 12.5|
| Employed                                | 65 | 38.7|
| Unemployed                              | 57 | 33.9|
| Retired                                 | 25 | 14.9|
| Familial psychiatric anamnesis          |    |     |
| Positive                                | 70 | 41.7|
| **Clinical variables**                  |    |     |
| Diagnosis                               |    |     |
| Depressive disorders                    | 54 | 32.1|
| Bipolar and related disorders           | 44 | 26.2|
| Schizophrenia spectrum and other psychotic disorders | 54 | 32.1 |
| Anxiety disorders                       | 5  | 3.0 |
| Obsessive–compulsive and related disorders | 11 | 6.5 |
| FRS                                     |    |     |
| Actual                                  | 55 | 32.7|
| Past                                    | 80 | 47.6|
| Lifetime                                | 86 | 51.2|

**FRS first rank symptoms**

### Discussion

The present study showed that ASI values were higher in patients with FRS, but they did not differ in patients with any specific psychiatric diagnosis. This outcome seemed reasonable, hypothesizing that ASI can vary depending on the current or lifetime presence of psychotic experiences. Furthermore, in the present sample of psychiatric patients, none of the diagnoses seemed to be related to the ASI score. Lelli describes ASI as a scale permitting to discriminate patients from controls, and patients with psychotic symptoms, from patients without them [26], demonstrating its validity and ability to individuate psychotic patients. As in other studies [28], the present study outlined not only the presence of FRS at study assessment but also as past or lifetime, to evaluate whether aberrant salience can be considered mainly a trait feature instead of a state characteristic, independently from the state of illness. Furthermore, the observation of the present...
confirmed the findings of previous studies, according to which FRS are not specific enough to provide a satisfactory diagnosis [29]. Although these analyses were conducted in a relatively small clinical sample, the results of the correlation analysis confirmed the reliability and consistency of collected data. Thus, the observed outcomes could provide interesting perspectives regarding the use of aberrant salience as a clinically useful paradigm, as part of a psychopathological approach evaluating vulnerability to psychosis recurrence. In this sense, higher values of ASI in patients with current, lifetime, and previous FRS seem to confirm this approach.

As the average ASI score in patients with a main diagnosis of schizophrenia did not significantly differ from the one observed in other psychiatric patients \( p = 0.107 \), possibly due to the small sample size, it is possible to hypothesize that salience alteration, underpinned by a
dysregulated dopaminergic firing, can be linked with a dimensional and trans-nosographic feature (highlighted by the correlation with FRS), rather than strictly fit into a diagnostic categorization.

According to Howes and Kapur’s III version, “A much more likely scenario is that a biological dysfunction may contribute to one of the major dimensions of the illness. The dopamine dysfunction is present even in subjects reflecting the extended phenotype—family members of people with schizotypy and symptomatic individuals at high risk of psychosis. Thus, the current evidence is consistent with dopamine hyperfunction being most closely linked to the dimension of psychosis […] Dopamine elevation appears specifically related more generally to psychosis proneness and not just to psychosis in schizophrenia” [30].

Considering both Kapur’s hypothesis concerning dopaminergic dysregulation underpinning salience alteration, and Howes’ theorization on altered dopaminergic release as the neurobiological basis of prodromal phases of psychosis [31], we suggest that ASI may be used as a tool to identify prodromal stages of psychosis.

The lack of correlation between ASI and psychiatric scales observed in this study may contribute to confirm its trans-nosographic clinical value. As aberrant salience is described as a part of primary psychotic experiences, this might be a more mindful and reasonable target of a neuroscientific investigation to overcome the established DSM and ICD classifications [32, 33]. The fact that this study was conducted in a single center, where only expert psychiatrists homogeneously performed assessments, reliability as a predictor of the psychotic disorder needs to be confirmed in a properly designed prospective studies, performed by consistently trained psychiatrists.

Conclusions
The results of this preliminary study seem to confirm ASI validity in identifying psychotic vulnerability. ASI may be used in screening and prevention programs to identify subjects for which a deeper and more careful psychopathological assessment is warranted. Aberrant salience conceptualization [34] may give a different framework in the assessment of psychotic diagnoses, underlining the importance of a dimensional approach in evaluating psychotic patients, through a trans-nosographic perspective. ASI

Abbreviations
AMDP: Manual for the Assessment and Documentation of Psychopathology; ASI: Aberrant Salience Inventory; CHAID: Chi-square automatic interaction detection; DSM: Diagnostic and Statistical Manual; FRS: First rank symptoms; GCP: Good clinical practices; HAMA: Hamilton Anxiety Scale; ICD: International Classification of Diseases; MADRS: Montgomery–Asberg Depression Rating Scale; MRS: Mania Rating Scale; PANSS: Positive and Negative Syndrome Scale; PLEs: Psychotic-like experiences.

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Authors’ contributions
AB, MT, MS, VR: designed the study. MT, MS, PM, CA, CB, LM, TT, FP, GD: collected patient’s data. AR performed the statistical analyses and drafted the article. All authors contributed to the interpretation of the data, revised and approved the manuscript draft, and are accountable for all aspects of the work according to ICMJE guidelines. All authors read and approved the final manuscript.

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Availability of data and materials
Data are available on request due to privacy/ethical restrictions.

Declarations
Ethics approval and consent to participate
This was a retrospective study that did not need ethical approval. All patients attending the visits at our center provide their consent to have their data used for studies ensuring patient’s privacy and in accordance to the Declaration of Helsinki. The study protocol was approved by the Ethics Committee of the University of Florence. Before they were enrolled in the trial, all patients received an information leaflet, in Italian language, explaining exhaustively the protocol and its implications. Further explanations were given by the research team as needed. Before collecting personal data and evaluating clinical conditions, patients were asked to sign their written informed consent where they provided their consent to publish the results of the study, too.

Consent for publication
All authors of the manuscript have read and agree to its content to be published on Annals of General Psychiatry.

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

Author details
1 Psychiatry Unit, Department of Health Science, University of Florence, Largo Brambilla 3, 50134 Florence, Italy. 2 Department of Mental Health and Addictions, Central Tuscany NHS Trust, Florence, Italy.

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