Adjunctive agents to antipsychotics in schizophrenia: a systematic umbrella review and recommendations for amino acids, hormonal therapies and anti-inflammatory drugs

Guillaume Fond,1,2,3 Jasmina Mallet,2,4 Mathieu Urbach,2,5 Michael Eriksen Benros,6,7 Michael Berk,8 Martina Billec,9 Laurent Boyer,10,3 Christop U Correll,11,12,13,14 Michele Fornaro,10,9 Jayashri Kulkarni,15 Marion Leboyer,2,16 Pierre-Michel Llorca,2,17 David Misdrahi,2,18 Romain Rey,2,19 Franck Schürhoff,2,16 Marco Solmi,20,21,22,23 Iris E C Sommer,24 Stephen M Stahl,25 Baptiste Pignon,16 Fabrice Berna.

ABSTRACT

Question This umbrella review and guidelines aimed to provide evidence to support the rational choice of selected adjunctive therapies for schizophrenia.

Study selection and analysis Following the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) and World Federation of Societies of Biological Psychiatry (WFSBP)-grading recommendations, 63 randomised control trials (RCTs) (of which 4219 unique participants have completed the RCTs) and 29 meta-analyses were analysed.

Findings Provisional recommendations (WFSBP-grade 1) could be made for two molecules in augmentation to antipsychotics: (1) N-acetyl-cysteine (NAC, 1200–3600 mg/day, for >12 consecutive weeks) in improving negative symptoms, general psychopathology (positive and negative syndrome scale for schizophrenia (PANSS) general psychopathology factor (G)-G subscale), with the RCTs with the longer duration showing the most robust findings; (2) polysaturated fatty acids (3000 mg/day of eicosapentaenoic acid, for >12 weeks) in improving general psychopathology. Weaker recommendations (ie, WFSBP-grade 2) could be drawn for sarcosine (2 g/day) and minocycline (200–300 mg/day) for improving negative symptoms in chronic schizophrenia (not early schizophrenia), and NAC for improving positive symptoms and cognition. Weak recommendations are not ready for clinical practice. There is provisional evidence that oestrogens and raloxifene are effective in some patients, but further research is needed to determine their benefit/risk ratio.

Conclusions The results of this umbrella review should be interpreted with caution as the number of RCTs included in the meta-analyses was generally small and the effect sizes were weak or medium. For NAC, two RCTs with low risk of bias have provided conflicting results and the WFSBP-grade recommendation included also the results of meta-analyses. These drugs could be provisionally prescribed for patients for whom no other treatments have been effective, but they should be discontinued if they prove ineffective.

WHAT IS ALREADY KNOWN ON THIS TOPIC

⇒ Antipsychotics are not effective in all patients with schizophrenia and some drugs have shown effectiveness in meta-analyses, but with inconsistent results.

WHAT THIS STUDY ADDS

⇒ An international expert committee has analysed the 63 randomised controlled trials for amino acids, anti-inflammatory drugs and hormonal therapies and has concluded that N-acetyl-cysteine and polysaturated fatty acids show an excellent benefit/risk ratio and should be recommended in clinical practice. Preliminary evidence for sarcosine and minocycline calls for further research.

INTRODUCTION

Antipsychotics currently represent the cornerstone treatment for schizophrenia.1 This class of drugs has transformed the course of the disease, essentially by reducing the positive symptoms, the duration of acute episodes and the risk of relapse.2 All antipsychotics to a varying degree block the D2 receptors in the striatum.3 It was hypothesised that this mechanism of action was a universal target in schizophrenia.4 However, the dopamine hypothesis is as simplistic for schizophrenia as the serotonin hypothesis for depression. For example, it has greater therapeutic translational validity for positive than negative or cognitive symptoms.3 Additionally, non-dopaminergic agents such as trace amine-associated receptor 1 (TAAR1) agents...
show promise. Overall, 30% of patients with schizophrenia do not respond to one antipsychotic and only 40% respond to clozapine, the antipsychotic that is indicated in those showing resistance to two antipsychotics. In addition, schizophrenia is heterogeneous. Some clinical subpopulations have been identified as potential new targets for precision medicine interventions. Among them, first-episode psychosis/schizophrenia, women, patients with chronic peripheral inflammation and/or oxidative stress, and treatment-resistant schizophrenia were considered in subgroup analyses in order to mitigate the heterogeneity of the response to antipsychotics reported in these subgroups.

During the last two decades, new biological mechanisms acting on psychosis but not directly on dopamine or its receptors are in late-stage development, emerging as effective antipsychotics. These include the M1/M4 muscarinic agonist xanomeline plus the peripherally restricted anticholinergic tropisium and several other procholinergic drugs, as well as the TAAR1 agent ulotaront. In addition, multiple meta-analyses have been published reporting the effectiveness of agents added to antipsychotics in schizophrenia targeting clinical subgroups and/or these pathophysiological pathways. Adjunctive antidepressants, such as mirtazapine and lamotrigine, have shown effectiveness in improving the symptomatology of schizophrenia and are recommended in some clinical guidelines. The role of augmentation strategies with other psychotropic drugs is less clear. Beyond psychopharmacologic medications, some promising non-psychotropic repurposed medicines have also been investigated in meta-analyses. However, while meta-analyses aim to provide data for effectiveness and tolerance, they are often not designed to readily inform the clinical practice.

Different meta-analyses may yield inconsistent results according to their inclusion criteria or their representativeness as new evidence becomes available for quantitative synthesis over time. Also, some randomised controlled trials (RCTs) with various risks of bias can swing the results in favour or against one treatment. The benefit/risk ratio is often not explored. When there are only small size RCTs for one treatment, a meta-analysis may overestimate or underestimate a treatment effect. Unless controlled via subgroup analyses, studies of high and low quality or studies from heterogeneous samples may be mixed together yielding spurious results. Despite statistical approaches (eg, funnel plot analyses), publication bias can further affect the results of meta-analyses as it can affect the development of recommendations based on clinical trials. The translation of these results into clinical practice prompts a different methodology developed through the 2019 World Federation of Societies of Biological Psychiatry (WFSBP)-grading recommendations. A consensual method to synthesise the evidence in psychiatry was published in 2018 by the WFSBP. For example, the WFSBP and the Canadian Network for Mood and Anxiety Treatments (CANMAT) societies have recently published recommendations for using nutrients in severe mental disorders. In their recommendations, only N-acetyl-cysteine (NAC) was recommended for treating negative symptoms of schizophrenia, and polyunsaturated fatty acids (PUFAs) were not recommended. However, these guidelines did not use the 2019 WFSBP methodology, and some evidence suggests that these recommendations could be updated or tempered.

Therefore, this work aimed to synthesise the available best-quality evidence on selected adjunctive treatments, including amino acids, hormonal therapies and anti-inflammatory drugs given adjunctively to current antipsychotics in order to guide clinical practice for the management of schizophrenia for clinicians and to provide evidence-based data for stakeholders and public policy makers.

**METHODS**

The detailed methods for literature search, inclusion criteria, data extraction, subgroups, risk of bias assessment and grading

---

**Table 1** Included randomised controlled trials (RCTs) with sample sizes

<table>
<thead>
<tr>
<th>Drug</th>
<th>RCTs (N)</th>
<th>RCTs with low/moderate/high risk of bias (N)</th>
<th>Total (N)</th>
<th>Drug (N)</th>
<th>Placebo (N)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N-acetyl-cysteine</td>
<td>8</td>
<td>3/4/1</td>
<td>523</td>
<td>258</td>
<td>265</td>
</tr>
<tr>
<td>Sarcosine</td>
<td>6</td>
<td>0/4/2</td>
<td>211</td>
<td>104</td>
<td>107</td>
</tr>
<tr>
<td>Minocycline</td>
<td>8</td>
<td>4/3/1</td>
<td>583</td>
<td>298</td>
<td>285</td>
</tr>
<tr>
<td>PUFAs</td>
<td>14</td>
<td>5/5/4</td>
<td>809</td>
<td>432</td>
<td>377</td>
</tr>
<tr>
<td>Oestrogens</td>
<td>9</td>
<td>5/3/1</td>
<td>677</td>
<td>383</td>
<td>294</td>
</tr>
<tr>
<td>Selective estrogen receptor modulator (raloxifene)</td>
<td>9</td>
<td>6/3/0</td>
<td>552</td>
<td>275</td>
<td>277</td>
</tr>
<tr>
<td>Aspirin</td>
<td>4</td>
<td>0/3/1</td>
<td>424</td>
<td>221</td>
<td>203</td>
</tr>
<tr>
<td>Celecoxib</td>
<td>5</td>
<td>1/3/1</td>
<td>440</td>
<td>222</td>
<td>218</td>
</tr>
<tr>
<td>Total</td>
<td>63</td>
<td>24/28/11</td>
<td>4219</td>
<td>2193</td>
<td>2026</td>
</tr>
</tbody>
</table>

PUFA, polyunsaturated fatty acid; SERM, selective estrogen receptor modulator.

---

**Figure 1** Forest plots of the main effects size of the selected adjunctive agents on total psychopathology. SERM selective estrogen receptor modulator. NAC, N-acetyl-cysteine; PUFAs, polyunsaturated fatty acids; SMD, standardised mean difference.
process are presented in online supplemental materials 1–7. The present review did not any receive financial support.

RESULTS
A total of 63 RCTs (4219 patients) and 29 meta-analyses were identified (presented in table 1 and online supplemental material 2). The detailed characteristics of the RCTs, efficacy results, risk of bias and Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) checklist are presented in online supplemental materials 3–5. The main effects size of the selected adjunctive agents on total psychopathology is presented in the forest plot of figure 1. The PRISMA flow chart is presented in figure 2. The detailed WFSBP-grade recommendations are presented in online supplemental materials 6 and 7. Importantly, we have changed the wording from ‘strong recommendation’ (corresponding to the GRADE 1 level) to ‘strong provisional recommendation’ to indicate that further RCTs may potentially modify the present recommendations (hence, provisional also applies to moderate and weak recommendations). The influence of sample size, risk of bias, patients’ groups, high-income versus upper middle-income countries are presented in online supplemental materials 8 and 9. We strongly encourage readers to carefully consider the materials on which the present recommendations are based.

DISCUSSION
Based on the present umbrella review, we found sufficient evidence to formulate strong provisional recommendations (WFSBP-grade 1) for three adjunctive agents when used in augmentation with antipsychotics in schizophrenia: (1) NAC (when used between 1200 and 3600 mg/day, for at least 12 weeks) with significant improvement in negative symptoms, general psychopathology and cognition, with the strongest evidence for the studies going at least 6 months; (2) PUFAs (with doses of 3000 mg/day of eicosapentaenoic acid (EPA), for at least 12 weeks) with significant improvement in general psychopathology; (3) transdermal estradiol (with doses of 0.1–0.2 mg/day) with significant improvement in positive symptoms and general psychopathology in childbearing aged women (especially ≥38 years old). In the latter case, data are, however, limited to 8-week trials. Recommendations regarding NAC and PUFAs are supported by meta-analyses concluding to a significant improvement without publication bias. However, a strong heterogeneity was also reported (67% for NAC and 38% for PUFAs, see figure 1). Weaker recommendations (ie, WFSBP-grade 2) could be made for other adjunctive agents when used in augmentation with antipsychotics for improving (1) negative symptoms: sarcosine (2 g/day), minocycline (200–300 mg/day), oestrogens (either daily 2 mg estradiol valerate or 0.625 mg conjugated oestrogens with 2.5 mg medroxyprogesterone acetate); (2) positive symptoms: NAC 1200 mg/day for at least 12 weeks or (3) general psychopathology: 2 mg estradiol valerate in men but only evaluated for 2 weeks.

The present work adds important findings to the previously published meta-analyses. In the light of these results, patients with respective negative, cognitive or general psychopathological symptoms could be encouraged to take NAC and/or PUFAs, as these agents are available over-the-counter. The findings pertaining to NAC are particularly intriguing, as they demonstrate moderate-term effects within the 12-week to 24-week timeframe. This raises important questions about the timing of observation and suggests that long-term assessments may yield additional valuable insights. Of note, in the case of NAC, the level Of Evidence (LoE) is only B but the grade recommendation was 1 due to its excellent acceptability, as recommended in the WFSBP guidelines. However, the off-label prescription of oestrogens and raloxifene in all women with schizophrenia cannot be recommended for sure due to safety issues (even in those aged >38 years in whom oestrogens and raloxifene seem more effective). The present results only confirm the efficacy of these agents that may still be an option in some case of resistance to conventional treatments.

In patients with an acute episode of schizophrenia, no adjunctive treatment in co-initiation to antipsychotics could be recommended with WFSBP-grade 1 evidence due to methodological issues of combining the augmentation of adjunctive agent with varying dosages of antipsychotics during the acute stabilisation phase. Yet, weaker recommendations (WFSBP-grade 2) could be made for improving (1) negative symptoms: sarcosine (2 g/day), minocycline (200–300 mg/day), raloxifene (120 mg/day) in men; (2) positive symptoms: PUFAs (at least 2000 mg/day EPA) for patients with low PUFAs levels, celecoxib (400 mg/day) or (3) general psychopathology: sarcosine (2 g/day), celecoxib (400 mg/day), raloxifene (120 mg/day) in men.

These recommendations are derived from the careful analysis of RCTs and meta-analyses representing the existing literature. To avoid any misleading interpretation of current practice, we would like to stress that the use of adjunctive treatment should come in second line after all attempts to optimise patients’ current antipsychotic treatment and psychosocial therapies have been made (dose optimisation, antipsychotic plasma level monitoring, managing comorbidity such as substance abuse and ruling out of somatic causes for non-response, etc) according to current recommendations. It is also important to carefully consider the benefit/risk balance before prescribing any adjunctive treatment. This is crucial to avoid augmentation with ineffective agents that carry a risk of side effects (including more severe negative symptoms). For instance, reducing the dosage of antipsychotics to their minimal effective dose may be safe (some studies suggest that it is not associated with a significantly increased risk of rehospitalisation compared with maintaining the treatment to the same dosage) and it may be an efficient
strategy to improve negative symptoms and cognition. This strategy should thus be first discussed with patients who stabilise after a first or multiple episode(s) before envisioning the use of adjunctive treatments for negative symptoms or cognition. There is consistent evidence for oestrogen augmentation in women with schizophrenia especially those over 38 years, but clinical implementation has not yet become common practice. The use of oestrogen needs to be done safely and concordant with existing practice guidelines, for gonadal hormone therapies. Oestrogen can be prescribed clinically as combined oral contraceptives for pre-menopausal women or through hormonal replacement therapy (estradiol patches, with regular progesterone addition) for post-menopausal women. Raloxifene could be an alternative for post-menopausal women. Further clinical research is required to determine the efficacy and safety for the clinical use of oestrogen therapies in the treatment of women with schizophrenia.

A somewhat paradoxical aspect of the present results should be mentioned regarding adjunctive treatment with the selected agents. The adjunctive therapies reviewed here were thought to target (1) particular biological pathways putatively involved in the pathophysiology of schizophrenia or (2) particular patients with schizophrenia presenting alterations of one of those biological pathways. However, almost all studies considered the schizophrenia group as a whole, and only a few examined or stratified for the kind of patients who were most likely to benefit from the tested drug. For example, although lower PUFA blood levels have been shown in schizophrenia as a group, only one RCT with a low risk of bias evaluating the effect of PUFA in schizophrenia took the blood level deficiency of PUFAs into account and showed significant improvement of positive symptoms only in this group. Similarly, among all RCTs testing the use of anti-inflammatory drugs, only one made the distinction between patients with/without low-grade peripheral inflammation (defined by CRP blood level ≥ 1 mg/L). Also, few studies testing the effect of adjunctive hormonal therapies tested hormone levels in the included patients (e.g., Kulkarni et al). Similarly, the interpretation of the results for general psychopathology is complex from a precision psychiatry perspective, as the PANSS-G factor encompasses heterogeneous symptoms such as anxiety, depression, lack of insight and attention disorders. All in all, these findings underscore the pressing need to validate the application of precision psychiatry approaches in future research. However, it is important to acknowledge that this also presents additional challenges in terms of study recruitment and feasibility.

Our results also suggest the benefits of applying the WFSBP-grade recommendations, as our conclusions differ somewhat from those of other meta-analyses. Our findings demonstrate that weak or moderate mean effect sizes in meta-analyses should not be directly translated into recommendations for or against prescribing a specific agent or group of agents. In fact, these effect sizes may correspond to a weak or limited level of evidence. Furthermore, additional complexities arise in trials comparing augmentation and co-initiation approaches. The discrepancies between the results of augmentation and co-initiation trials indicate the relevance of this distinction. In co-initiation trials, the control groups receive an active antipsychotic treatment that reduces psychotic symptoms, while augmentation trials typically involve a stable antipsychotic treatment in the control group, which may result in potentially weaker and clinically significant variations. However, it is important to highlight certain features of the WFSBP-grade system to clarify our recommendations. The designation of ‘limited evidence’ (WFSBP-grade 2) can be assigned when the agents are well tolerated and when one of the following conditions is met: (1) a single RCT with low or moderate risk of bias shows significant improvement without other RCTs of equal quality demonstrating non-significant results, or (2) two RCTs with low risk of bias yield contradictory results, but a meta-analysis demonstrates significant improvement. Therefore, the conclusions may be influenced by both the number of RCTs and the quality of the evidence. This findings should be taken into consideration when interpreting the recommendations for certain adjunctive treatments such as sarcosine, minocycline or oestrogens, as there were generally few RCTs available. It also emphasises the need for new large-scale studies of high quality and low risk of bias to confirm the validity of our recommendations.

It is important to acknowledge that, despite following the SIGN recommendations for grading the risk of bias, there remains a potential for global subjectivity (as mentioned in the SIGN method) regarding the final level of evidence. From this point of view, our grading may appear more stringent compared with the results of some meta-analyses, but it must be acknowledged on the contrary that other guidelines such as the NICE or the GRADE system have a more stringent and may thus come to distinct recommendations than ours. Given that our objective was to support evidence-based practice, we aimed to provide the most rigorous recommendations without dismissing potentially effective agents. Additionally, the risk of bias is also supported by treatment allocation concealment. Blinding between active agents and placebo is not always straightforward, as some active agents may induce noticeable adverse events, which could compromise blinding. This may not be the case for amino acids but may be more common for other active agents included in our work. For example, minocycline could cause diarrhoea, and aspirin may result in easy bruising, which may indicate to participants that they are receiving the active agent. Furthermore, it was not always clear in trials involving PUFAs whether the placebo was comparable with PUFA treatment in terms of flavour. It is worth noting that the same criticism can be raised regarding the side effects of common antipsychotics.

This area of research has several limitations that should be acknowledged. First, we focused this review on augmentation strategies of antipsychotics in schizophrenia, as psychotropic augmentation strategies of antipsychotics have been comprehensively reviewed in a previous umbrella review. Second, the number of RCTs included in the meta-analyses assessed in this umbrella review was generally small so that the inclusion of data from new low-risk-of-bias randomised controlled trials carries a high likelihood of potentially influencing the current recommendations in one way or another. Consequently, we have opted to make ‘provisional’ recommendations for some molecules rather than definitive recommendations. To mitigate this limitation as much as possible, we conducted additional literature searches and included any newly published RCTs since the completion of the last meta-analysis. Third, there was heterogeneity in study designs, populations, study durations and intervention doses. To address this relevant heterogeneity, particularly in terms of illness phase and augmentation versus co-initiation strategies, we stratified the results and recommendations based on these important factors. It is important to note that our recommendations were not based on the effect size of each drug, as individual patient responses can vary greatly. Additionally, meta-analyses yielded different effect sizes due to the inclusion of RCTs with a high risk of bias. The focus of our recommendations was solely on the superiority of the add-on strategy with an active agent compared with add-on placebo. While there may have been a
potential bias in low sample size studies regarding cognition, we found no evidence of such bias for positive and negative symptoms. The fact that low risk of bias studies had a higher likelihood of showing significant effects on general psychopathology further supports the validity of our results.

Fourth, we also found that the occurrence of positive significant results for negative symptoms was more frequent in studies conducted in upper-middle-income countries compared with high-income countries (see online supplemental material 9). This difference was consistently observed when analysing studies with low risk of bias or specifically focusing on NAC, oestrogens and PUFAs. Therefore, one could argue that our recommendations for these drugs may have been biased by those particular studies. This also justifies our use of the term ‘provisional’, as further studies conducted in high-income countries may help confirm our recommendations. Additionally, numerous findings have been provided by Chinese and Iranian research teams, and some authors have shown that accurately assessing the bias risk of these studies can be challenging and can possibly hide high risk of bias. The inclusion of Chinese and Iranian researchers in future recommendations could help address this issue.

Fifth, the field of adjunctive treatments is more complex to explore and analyse than antipsychotic monotherapy for multiple reasons exposed in our results. While almost all low risk of bias RCTs used fixed doses, an important number of high risk of bias RCTs did not adequately describe the antipsychotics administered in each arm at baseline and during the trial or did not check if these treatments were comparable between groups at the end of the trial. In these RCTs, the described effect could be due to a significant effect of the adjunctive drug or to a change of antipsychotic dose in one of the arms. WFSPB guidelines ensured that no recommendation was influenced on this kind of bias by downgrading the level of evidence when necessary. In addition, for several adjunctive drugs, the evidence was based on many small studies. However, as the overall meta-regression analyses from the study by Jeppesen et al showed a decreasing effect with the sample size, large low risk of bias studies are needed to confirm our recommendations, thus keeping in mind that sample size had no relevant influence on the probability to find a significant effect on the 63 studies included (see online supplemental material 9). Sixth, while one strength of our systematic umbrella review and recommendations is that we made the distinction between some symptom dimensions of schizophrenia, this may also account for some differences between our recommendations and previous ones, for instance, regarding PUFAs. However, we found that no data to evaluate functioning and patient-important outcomes and other important information on costs or adverse events (including those like irritability or weight gain) were not or not fully assessed due to limited data availability. The present recommendations are therefore only based on clinician-rated symptoms of the PANSS (for most studies) and the effect on quality of life and daily-life functioning is unclear. Given that patient input is crucial, particularly for weak recommendations where the benefit/risk balance is uncertain, future recommendations should include patients.

Seventh, due to lack of data, we were not able to integrate depression or anxiety in our outcomes. Depression is present in at least one-third of patients with schizophrenia, and has a major impact on prognosis and quality of life. Including depression (optimally with the specific Calgary Depression Rating Scale) is strongly recommended for future research and might be useful to appreciate a putative antidepressant effect of PUFAs in schizophrenia, as previously suggested in mood disorders. Finally, since the search for the study ended in February 2022, it potentially missed more recent evidence that will need to be included in future, up-to-date studies.

**CONCLUSION**

Based on the existing literature, the adjunctive use of NAC and PUFAs can be tentatively recommended in all phases of schizophrenia, considering their potential benefits for negative symptoms and/or general psychopathology, as well as their high acceptability. NAC also appears to be the only adjunctive treatment that may have potential benefits for cognition. However, it should be emphasised that these recommendations are solely based on a limited number of RCTs, and the current recommendations are derived from significant results and risk of bias rather than effect sizes. These drugs should potentially be prescribed for patients for whom antipsychotics have proven ineffective in addressing negative symptoms and/or general psychopathology and/or cognitive function. Furthermore, if these drugs are found to be ineffective, they should be discontinued. The term ‘provisional’ has been used to underscore this point. Transdermal estradiol, used for short-term, can also be provisionally recommended in women of childbearing age to improve positive symptoms. Other adjunctive agents, such as sarcosine, minocycline, oestrogens or SERMs, may also be effective in certain clinical scenarios, but additional low-risk RCTs are needed. It is important to note that some safety concerns should be considered for these agents, with the exception of sarcosine. These recommendations are provisional and emphasise the need for further research on targeted approaches, such as selecting or stratifying patients based on inflammatory markers or nutrient levels. Large-scale studies with low risk of bias are required to identify the patients who are most likely to benefit from a specific adjunctive agent.

**Author affiliations**

1Department of psychiatry, Assistance Publique des Hôpitaux de Marseille, Marseille, France
2Fondation FondaMental, Creteil, France
3CERES-Health Service Research and Quality of Life Center, AMI, Marseille, France
4Department of Psychiatry, Louis Mourier Hospital, Colombes, France
5Department of Adult Psychiatry and Addictology, Centre Hospitalier de Versailles, Le Chesnay, France
6Copenhagen Research Center for Biological and Precision Psychiatry, Mental Health Centre Copenhagen, Copenhagen, Denmark
7Department of Clinical Medicine, Faculty of Health and Medical Sciences, University of Copenhagen, Copenhagen, Denmark
8Deakin University, School of Medicine, and Barwon Health; IMPACT, the Institute for Mental and Physical Health and Clinical Translation; Orygen The National Centre of Excellence in Youth Mental Health, The Florey Institute of Neuroscience and Mental Health and the Department of Psychiatry, University of Melbourne and the Department of Public Health and Preventive Medicine, Monash University, Melbourne, Victoria, Australia
9Department of Neuroscience, Reproductive Sciences, and Dentistry, Section of Psychiatry, Federico II University of Naples, Naples, Italy
10Département d’information médicale, Assistance Publique des Hôpitaux de Marseille, Marseille, France
11Department of Psychiatry, Zucker Hillside Hospital, Glen Oaks, New York, USA
12Department of Psychiatry and Molecular Medicine, Donald and Barbara Zucker School of Medicine at Hofstra/Northwell, Hempstead, NY, USA
13Department of Child and Adolescent Psychiatry, Charité Universitätsmedizin Berlin, Berlin, Germany
14German Center for Mental Health (DZPF), partner site Berlin, Germany
15Department of Psychiatry, Monash Alfred Psychiatry Research Centre, Alfred Hospital and Monash University Central Clinical School, Monash University,607StKildaRd, Level4, Melbourne, Victoria, Australia 3004, Melbourne, Victoria, Australia
16Department of Psychiatry, Univ Paris-Est-Créteil (UPEC), AP-Hôpitaux Universitaires « H. Mondor », DMU IMPACT, INSERM U955, INRHB, translational Neuropsychiatry, F-94010 Créteil, France, Créteil, France
17Département de psychiatrie, Université Clermont Auvergne, CMP-B CHU, CNRS, Clermont Auvergne INP, Institut Pascal, Clermont-Ferrand, France
18Department of psychiatry, Univ Paris-Sud - AP-HP, Hôpitaux Universitaires « H. Mondor », DMU IMPACT, INSERM U955, INRHB, translational Neuropsychiatry, F-94010 Créteil, France, Créteil, France
19Département de psychiatrie, Université Clermont Auvergne, CMP-B CHU, CNRS, Clermont Auvergne INP, Institut Pascal, Clermont-Ferrand, France

open access

In 6 Fond G, et al. BMI Ment Health 2023;26:1–7. doi:10.1136/bmjment-2023-300771

REFERENCES

7. Mustafa FA. Could some patients with treatment-resistant schizophrenia have reversible conditions. Psychiotor Psychosom 2018;87:369.
21. Tsai GE, Lin P-Y. Strategies to enhance N-methyl-D-aspartate receptor-mediated neurotransmission in schizophrenia, a critical review and meta-analysis. Singh SP, Singh V. Meta-analysis of the efficacy of adjunctive NMDA receptor modulators in chronic schizophrenia. CNS Drugs 2011;25:859–85.