PREVENTIVE TREATMENTS FOR PSYCHOSIS: UMBRELLA REVIEW
(JUST THE EVIDENCE)

– Systematic review –

Running title: Prevention of psychosis – just the evidence

Paolo Fusar-Poli1,2,3,4, Cathy Davies1, Marco Solmi1,5, Natascia Brondino3, Andrea De Micheli1, Magdalena Kotlicka-Antczak6, Jae Il Shin7, Joaquim Radua1,8,9*.

Affiliations

1Early Psychosis: Interventions and Clinical-detection (EPIC) lab, Department of Psychosis Studies, Institute of Psychiatry, Psychology & Neuroscience, King's College London, London, UK;
2OASIS service, South London and Maudsley NHS Foundation Trust, London, UK;
3Department of Brain and Behavioral Sciences, University of Pavia, Pavia, Italy;
4National Institute for Health Research, Maudsley Biomedical Research Centre, South London and Maudsley NHS Foundation Trust, London, UK;
5Neuroscience Department, Psychiatry Unit, University of Padua, Padua, Italy;
6Department of Affective and Psychotic Disorders, Medical University of Lodz, Lodz, Poland;
7Department of Pediatrics, Yonsei University College of Medicine, Seoul, Republic of Korea;
8Institut d'Investigacions Biomèdiques August Pi i Sunyer (IDIBAPS), CIBERSAM, Barcelona, Spain;
9Department of Clinical Neuroscience, Centre for Psychiatric Research and Education, Karolinska Institutet, Stockholm, Sweden.

* Joint last authors.

Abstract: 349
Text: 9542
Tables: 5
Figures: 5
Endnote reference file: Prognosis.enl

Correspondence to: Dr Paolo Fusar-Poli MD PhD, Department of Psychosis Studies, 5th Floor, Institute of Psychiatry, Psychology & Neuroscience, PO63, 16 De Crespigny Park, SE5 8AF London, UK. E-mail: paolo.fusar-poli@kcl.ac.uk
ABSTRACT

Background
Indicated primary prevention in young people at Clinical High Risk for Psychosis (CHR-P) is a promising avenue for improving outcomes of the most severe mental disorders; their effectiveness has recently been questioned.

Methods
Umbrella review. A multi-step independent literature search of Web of Science until January 11th, 2019 identified interventional meta-analyses in CHR-P individuals. The individual RCTs that were analysed by the meta-analyses were extracted. A review of ongoing trials and a simulation of living meta-analysis complemented the analysis.

Results
Seven meta-analyses investigating preventive treatments in CHR-P individuals were included. None of them produced pooled effect sizes across psychological, pharmacological or other types of interventions. The outcomes analysed encompassed: risk of psychosis onset, the acceptability of treatments, the severity of attenuated positive/negative psychotic symptoms, depression, symptom-related distress, social functioning, general functioning, and quality of life. These meta-analyses were based on 20 RCTs: the vast majority defined the prevention of psychosis onset as their primary outcome of interest and only powered to large effect sizes. There was no evidence to favour any preventive intervention over any other (or control condition) for improving any of these clinical outcomes. Caution is required when making clinical recommendations for the prevention of psychosis in CHR-P individuals.

Discussion
Prevention of psychosis from a CHR-P state has been and should remain, the primary outcome of interventional research, refined and complemented by other clinically meaningful outcomes (severity of symptoms, functional status, recovery, quality of life). Stagnation of knowledge should promote innovative and collaborative research efforts, in line with the progressive and incremental nature of medical knowledge. Advancements will most likely be associated with the development of new experimental therapeutics that are ongoing, along with the ability to deconstruct the high CHR-P heterogeneity. This would require the estimation of treatment-specific effect sizes through living individual participant data meta-analyses, controlling risk enrichment during recruitment, statistical power and embedding precision medicine.
within youth mental health services that can accommodate sequential prognosis and advanced trial designs.

**Conclusions**
The evidence-based challenges and proposed solutions addressed by this umbrella review can inform the next generation of research into preventive treatments for psychosis.

**Keywords:** Psychosis, schizophrenia, prevention, treatments, meta-analysis, evidence
INTRODUCTION

Prevention of psychosis is a promising avenue for improving outcomes of one of the most severe mental disorders[1]. It entails three stepped core components: efficient detection of individuals at risk, an accurate prognosis of outcomes and an effective preventive treatment that can impact the course of the disorder[2]. These activities are usually managed by specialised clinical services. The first rate-limiting step is the detection of children, adolescents and young adults aged 8-40[3] (more frequently 14-35)[4] who may be at risk of developing psychosis. Their detection is based on recruitment campaigns[5] that filter individuals who have accumulated several risk factors[6] for the development of psychosis, thus enriching the level of risk. The second step is the clinical assessment of these individuals and the formulation of a prognosis[7; 8; 9]. Individuals meeting psychometric intake criteria for a Clinical High Risk state for Psychosis (CHR-P[10]) are functionally impaired[11] and have a 20% risk of developing psychosis at 2 years[12] (but not an increased risk of developing other non-psychotic mental disorders[13; 14]). The third, final step is the provision of an effective intervention, which can impact the clinical outcomes of CHR-P individuals. Evidence of effective interventions in CHR-P individuals—also termed as primary indicated interventions[15]—will be the focus of the current manuscript.

We first review the historical development of preventive treatments for psychosis by adapting the “Gartner Hype Cycle[16]” model to the CHR-P paradigm (for details see[17]). The Hype Cycle has been used to describe the course of knowledge in other areas of medical knowledge (e.g. machine-learning[18]). The Gartner Hype Cycle describes the course of new technological discoveries, assuming that humans tend to overestimate the impact of a new discovery in the short term, while largely underestimating the same in the long term. The Hype Cycle—illustrated in Figure 1—includes 5 different stages: (1) innovative trigger, (2) peak of inflated expectations, (3) trough of disillusionment, (4) slope of enlightenment, and (5) plateau of productivity or knowledge.

With reference to the CHR-P paradigm, the innovative trigger for the development of preventive treatments for psychosis was the set-up of the first CHR-P clinic (in 1995), followed (in 2002) by the publication of the first randomised controlled trial in CHR-P individuals. Such a trial employed antipsychotic (risperidone) and psychological
treatment, and demonstrated that they were effective in reducing the risk of developing a first episode of psychosis in CHR-P samples[19]. Over the ensuing years, the innovative trigger led to an explosion of enthusiasm with the development of additional treatments that—it was hoped—could prevent the onset of psychosis from a CHR-P state. Randomised controlled trials involving antipsychotics (olanzapine, 2006[20]), psychological therapies (Cognitive Behavioural Therapy[21], 2004 or Integrated Psychological Intervention[22], 2012) and dietary interventions (Omega-3[23]) all confirmed some degree of efficacy for reducing the onset of psychosis in CHR-P individuals (Figure 1). The peak of inflated expectations and optimism was likely reached through the publication of the National Institute for Health and Care Excellence (NICE) Clinical Guidelines 178 (2014)[24] and the European Psychiatric Association (EPA) guidelines (2015[25]). These guidelines set the gold standard for the prevention of psychosis in CHR-P individuals in clinical routine. At the same time, these guidelines also represented the starting point of the trough of disillusionment, because they were grounded on "non-conclusive[26]" and "moderate quality[26]" evidence of small magnitude (risk difference −0.07: 95%CIs −0.14 to −0.01)[26]. Upon closer inspection, the two guidelines were already reflecting a lack of robust evidence at the time of their release, because they were partially discordant. In fact, while prophylactic treatment with antipsychotics was altogether not recommended by NICE guidelines[24], the EPA allowed their use in the case of severe and progressive symptomatology[24]. Over the following years, the trough of disillusionment for effective preventive treatments for psychosis onset received additional corroborating support. Two large randomised controlled trials showed that neither Cognitive Behavioural Therapy (2012[27]) nor Omega-3 interventions (2017/2018[28; 29]) are effective in reducing the progression to psychosis from a CHR-P state. Overall, seven new trials involving 992 new CHR-P participants (an increase of more than 50%) have been published since the last meta-analysis that informed the NICE guidelines[30]. All of these trials were negative[30; 31]. An updated network meta-analysis (2018)[30] incorporating these new randomised controlled trials confirmed the lack of evidence to support specific preventive treatments over each—and every—other for the prevention of psychosis in CHR-P individuals.

The current trough of disillusionment phase has stimulated discussion across the field. On the one hand, some authors have attempted to minimise the clinical relevance of
the evidence itself. Specifically, it has been argued that current preventive treatments are actually effective on outcomes other than the prevention of psychosis onset, that recent network meta-analyses should have estimated effects across different types of preventive treatments “pooled together”, and that the primary outcome analysed by these meta-analyses—i.e. transition to psychosis from a CHR-P state—is a “second-order issue” of questionable clinical significance[32]. On the other side, some authors have argued that there is now enough evidence to abandon the CHR-P paradigm altogether. The current study explores the area between these two polarities, to recognise the key challenges while at the same time proposing ways to overcome them. To address such an uncertain phase of knowledge, we strictly adopt an evidence-based medicine approach. We will interrogate the published literature by conducting an umbrella review[33], i.e. a review of meta-analyses in the field of preventive interventions for psychosis in CHR-P individuals. Because umbrella reviews are based on meta-analyses previously published, they represent a bird’s-eye view on a determinate topic and reflect one of the highest levels of evidence. First, we will specifically review the evidence that preventive interventions can impact clinical outcomes (including and going beyond the development of psychosis) in CHR-P individuals. We will also review the use of pooled vs treatment-specific effect size estimates. Second, we will systematically review the extent to which prevention of psychosis has been the first or second-order outcome in interventional studies in this population. Third, we will critically interpret the umbrella review findings to inform the next slope of enlightenment stage, which will hopefully deliver a plateau of knowledge in this field.

METHODS
Search strategy, selection criteria and data extraction for the systematic review
A multi-step literature search was performed. First, systematic searches were conducted in the Web of Science (which includes Web of Science Core Collection, BIOSIS Citation Index, KCI - Korean Journal Database, MEDLINE, Russian Science Citation Index, and SciELO Citation Index), until January 11th, 2019, with no restrictions on language or publication date. The keywords “psychosis risk” or “clinical high risk” or “at risk mental state” or “prodromal”, “CAARMS” or “SIPS” or “basic symptoms” or “UHR” or “CHR” were used, filtering for the category “review” and “psychiatry” through the Web of Science categories function. Second, we searched
the abstracts of retrieved articles to include only those that employed meta-analytical methods. Third, full-text articles identified by this process were then screened and inspected against the inclusion and exclusion criteria. The literature search, study selection and data extraction were conducted by two authors independently. During all stages, in the case of disagreement, the consensus was reached through discussion with a third author.

Studies were eligible for inclusion when the following criteria were fulfilled: a) meta-analyses (aggregate or network) in CHR-P individuals (defined according to established international criteria[3]); b) a clear and primary focus on interventions for this patient population, with no restriction on the design of the primary studies (open-label, controlled/uncontrolled, randomised/unrandomised, naturalistic studies) or on the type of the intervention (medications, psychological, physical, dietary, experimental); c) investigating clinical outcomes in CHR-P individuals. The exclusion criteria were: a) original studies, study protocols, abstracts, systematic reviews without quantitative analyses and any other non-meta-analytical study; and b) lacking a clear primary focus on interventions for CHR individuals. In the case of two or more articles addressing the same outcome we selected individual participant data meta-analyses over aggregate network meta-analyses over pairwise-meta-analyses, and, in a second step, the most recent study. This was done to respect the hierarchy of the evidence.

Research is conducted at different levels: primary research consists of original studies while secondary research comprises qualitative reviews, systematic reviews, and meta-analyses. Within meta-analyses, network meta-analyses offer additional benefits over standard pairwise analyses in that the comparative efficacy of specific interventions can be estimated and ranked, even when two treatments have never been compared directly head-to-head. Furthermore, since network meta-analyses can improve the precision of estimates by allowing integration of both direct and indirect treatment effect estimates, it is recommended over pairwise meta-analyses by the World Health Organization as a basis for clinical guidelines. Therefore, network meta-analyses should be considered the highest level of evidence in CHR-P treatment guidelines. Finally, Individual Participant Data meta-analyses should be considered as a higher level of evidence compared to aggregate-data meta-analyses, because they can improve the quality of the data, the analyses and thus the reliability of the results. Therefore, they are considered the gold standard. The
quality of the meta-analyses identified through the literature search was assessed with the Assessing the Methodological Quality of Systematic Reviews (AMSTAR) tool[35]. AMSTAR is a reliable and valid measurement tool to assess the methodological quality of systematic reviews. AMSTAR assess whether the systematic review was based on an "a priori" protocol, if authors run screening and data extraction in duplicates, if they run a comprehensive search, including grey literature, if they provided a complete list of both included and excluded studies (at full-text assessment), if they assessed quality of included studies, and accounted for such quality in conclusions, if used appropriate methods to pool and analyse data, if assessed publication bias, and if disclosed information on potential conflict of interest. AMSTAR score goes from 0 (low quality) to 11 (high quality)[35].

In a subsequent step, we additionally extracted the individual studies that were analysed by the corresponding meta-analysis that was included in the current umbrella review. Specifically, we included only randomised controlled trials with no restriction on study design, being published in the English language and with a sample size of at least 10 participants[36]. Finally, to further provide a comprehensive view of current research in this field we systematically searched the clinicaltrials.gov using the keyword “psychosis risk” and filtering for randomised trials that were recruiting or active but not recruiting. The records were then summarised in a descriptive table.

**Simulation of a living meta-analysis**

Unfortunately, meta-analyses—even if based on individual participant data—are outdated as soon as new studies on the same topic emerge. Once published, only a minority of meta-analyses are then updated within two years of publication[37]. Such an inability to maintain recency may lead to significant inaccuracy in clinical practice which is not updated with evidence-based medicine. For example, by 2 years post-publication, 23% of non-updated meta-analyses will have failed to incorporate new evidence that would substantively change its conclusions[38].

Cumulative meta-analysis, defined as updating a meta-analysis whenever a new eligible RCT becomes available, can be used to address this[39]. However, the problem is that the median time taken for a primary study to be incorporated into a meta-analysis ranges from 2.5 to 6.5 years[40]. Thus, in 2014, “living systematic
reviews" were proposed as a framework for continuously updating meta-analyses[41] (Figure 2).

Living meta-analyses are particularly indicated when the question to be addressed is essential to decision-making, when there is some uncertainty of the evidence, when new information is likely to change the findings and when there is likely to be new evidence[42]. All of these conditions apply to the CHR-P field. Crucially, living meta-analyses require bespoke analytical approaches that have been developed for sequential (interim) analyses of randomised clinical trials[43; 44; 45; 46]. We simulate here a living meta-analysis using the available RCTs. We employed methods which allow reducing the chances of inflating type-I errors (by using the alpha-spending monitoring boundaries[47]) and type II errors (by estimating the required "a priori anticipated information size" (APIS), i.e., the minimum required sample size to detect an assumed minimal clinically important effect-size—as recommended by experts[48]—with a prespecified statistical power)[49]. Furthermore, we used approximate Bayesian approaches to reduce the misestimation of heterogeneity[50]. The results are presented in specific plots to facilitate the interpretation of the core findings.

RESULTS

Database for the systematic review

The initial literature search (Figure 3) retrieved 2328 records, of which only 327 were meta-analyses and were screened on the basis of title and abstract reading. Most of them were not focusing on CHR-P individuals or interventions, and only 14 full-text aggregate meta-analyses were eventually assessed for eligibility. There were no individual participant data meta-analyses.

One of these 14 meta-analyses was not reporting quantitative data[51] and 6 were overlapping with more recent meta-analyses[25; 52; 53; 54; 55; 56]: these studies were thus excluded. Since two network meta-analyses addressed the same outcome and were published in the same year (2018)[57; 58], it was not possible to apply our exclusion criteria. Both network meta-analyses were thus included. The final database comprised 7 meta-analyses: 4 aggregate-data network meta-analyses and 3 aggregate-data pairwise meta-analyses. Additionally, two errata were retrieved, but
they were not counted as independent articles[59; 60]. The full list of meta-analyses is listed in Table 1.

**Effect of preventive treatments on clinical outcomes in CHR individuals.**

*Prevention of psychosis onset*

An aggregate-data network meta-analysis published in 2018 showed a lack of evidence to favour specific preventive treatments compared to each—and every—other for the prevention of psychosis onset from a CHR state[61]. These comparisons also included needs-based interventions or placebo. Some authors speculated that these findings would have changed if the Neurapro trial[28], which was classified as Omega-3 + needs-based intervention vs needs-based intervention (in line with other independent network meta-analyses[58; 62]) was classified as Omega-3 + Cognitive Behavioural Therapy (Cognitive Behavioural Case Management) + needs-based interventions vs cognitive behavioural therapy + needs based interventions[32]. As already mentioned in the main paper[30], sensitivity analyses testing different definitions of the nodes did not change the results. For comprehensiveness, we further append in eFigure 1 the forest plot of the network meta-analysis when the Neurapro trial was coded as Omega 3 + Cognitive Behavioural Therapy (Cognitive Behavioural Case Management) + needs-based interventions vs cognitive behavioural therapy + needs-based interventions. The results remained unchanged.

*Acceptability of treatments*

An aggregate-data network meta-analysis published in 2018 showed a lack of evidence to favour any specific preventive treatment compared to every other (including also needs-based interventions or placebo) with respect to acceptability (defined as treatment drop-outs for any reason)[61].

*Symptoms*

Two independent aggregate-data network meta-analyses published in 2018 found lack of evidence to favour specific preventive treatments compared to each (and every) other for improving attenuated positive psychotic symptoms[58; 63]. Another aggregate-data network meta-analysis published in 2018 demonstrated no evidence to favour specific preventive treatments compared to each (and every) other
for the improvement of attenuated negative psychotic symptoms[62]. Some authors speculated that this meta-analysis showed “trend-level” benefits for N-methyl-D-aspartate receptor (NMDAR) modulators compared to placebo[32]. However, the 95%CI of the Standardised Mean Difference included the null hypothesis (from −1.09 to 0.02)[62]. The authors of the network meta-analysis in a following erratum clarified that neither efficacy nor effectiveness was statistically confirmed for any of the examined treatments and that “the abstract and text contained a misstatement regarding effectiveness”[60].

An aggregate-data pairwise meta-analysis published in 2013 indicated that preventive treatments have no impact on depression in CHR-P individuals[26].

A further aggregate-data pairwise meta-analysis published in 2014 found that preventive treatments have no impact on symptom-related distress in CHR-P individuals[64].

**Functioning**

An aggregate-data pairwise meta-analysis published in 2018 found no effect of available preventive treatments for the improvement of social functioning in CHR-P individuals[65]. An aggregate-data pairwise meta-analysis published in 2014 found no effect of preventive treatments on a broader level of functioning[64].

**Quality of life**

The most recent meta-analysis to explore the impact of preventive treatments on quality of life in CHR-P individuals was published in 2014. There were no significant treatment effects reported[64].

**Pooled vs specific-treatment effect sizes**

None of the included meta-analyses estimated overall effect size across different categories of treatments (i.e. medications, psychological, dietary) pooled together.

**Primary outcomes investigated by randomised controlled trials in CHR-P individuals**

As indicated in the methods, we further extracted details of the randomised controlled trials that were analysed by the meta-analyses included in the current umbrella review. A total of 20 trials were retrieved (Table 2). The treatments tested were:
• needs-based interventions;
• omega-3 + needs-based interventions;
• ziprasidone + needs-based interventions;
• olanzapine + needs-based interventions;
• aripiprazole + needs-based interventions;
• amisulpride + needs-based interventions;
• Integrated Psychological Interventions;
• family therapy + needs-based interventions;
• D-serine + needs-based interventions;
• Cognitive Behavioural Therapy, French & Morrison protocol + needs-based interventions;
• Cognitive Behavioural Therapy, French & Morrison protocol + risperidone + needs-based interventions;
• Cognitive Behavioural Therapy, van der Gaag protocol + Cognitive Behavioural Therapy, French & Morrison protocol + needs-based interventions;
• cognitive remediation therapy;
• computer games.

The primary outcome investigated by these trials as well as the rationale provided to support it are reported in Table 2. The vast majority (14/20, 70%) of randomised controlled trials defined the prevention of psychosis onset from a CHR-P state as their primary outcome. Three studies defined as primary outcome the reduction of attenuated psychotic symptoms\[66; 67; 68\], two studies the improvement of cognitive functioning\[69; 70\] and one study the improvement of social functioning\[71\]. The AMSTAR scoring for each meta-analysis included is reported in Table 2.

**Ongoing clinical trials identified in CHR-P individuals**

Ongoing experimental randomised trials listed in clinicaltrials.gov that are currently recruiting CHR-P patient are summarised in Table 3. Other experimental treatments that are not yet listed in the clinicaltrials.gov database are presented below.

**Oxytocin**
Impairments of social cognition, which include emotional processing, theory of mind and attributional style, are a primary cause of disability in psychosis and respond poorly to current treatments. Emotional dysregulation and social cognition problems are also common in CHR-P individuals, are a key source of distress and contribute to loss of functioning. Oxytocin, a neuropeptide, has numerous prosocial and antipsychotic-like effects in animals. In healthy individuals, oxytocin promotes interpersonal trust, social interactions and emotional bonding whilst decreasing arousal and aversion towards negative or threatening social stimuli (for review see). In patients with psychosis, oxytocin has been shown to improve emotional recognition and social dysfunction and may ameliorate psychotic symptoms, although the hope of its potential as a treatment in established psychosis has been somewhat stifled by recent negative meta-analytic findings. Nevertheless, neuroimaging studies demonstrate that oxytocin can modulate various indices of brain function, both task-specific and at rest, and in regions critically implicated in the onset of psychosis, such as limbic and midbrain brain regions. A randomised, double-blind, placebo-controlled, crossover acute-challenge MRI study has recently shown that oxytocin modulates hippocampal perfusion in CHR-P individuals, providing the first neurophysiological evidence for disease-target engagement. Further studies are ongoing that will better clarify the neurobiological effects of oxytocin in this patient population.

Cannabidiol

Cannabidiol is a major constituent of cannabis. In contrast to the psychoactive cannabinoid Δ9-tetrahydrocannabinol (THC), the non-psychoactive compound cannabidiol shows anxiolytic and potential antipsychotic properties. Early findings that administration of cannabidiol reduced psychotic symptoms in patients with established psychosis led to the suggestion that it may also have therapeutic potential in those at CHR-P. Interest in cannabidiol is enhanced by its unique mechanism of action compared to established antipsychotics, as well as its distinct lack of serious adverse effects. A randomised controlled trial using cannabidiol and MRI in CHR-P individuals has recently been completed. In 33 CHR-P individuals and 19 healthy controls, cannabidiol was found to modulate activation of brain regions strongly implicated in the onset of psychosis during verbal learning, such as the striatum, medial temporal cortex and midbrain. These early results support the view that cannabidiol may be
an effective treatment strategy. Future large-scale randomised controlled trials involving cannabidiol are expected from different research centres worldwide.

Simulation of the living meta-analysis

The results of the simulated living meta-analyses is provided in Figures 4 and 5.

Figure 4 illustrates the theoretical results of a cumulative living meta-analysis investigating the efficacy of CBT versus needs-based intervention on the risk of developing psychosis at 6-months, for a treatment which can halve the risk of psychosis (Risk Ratio Reduction, RRR=0.5). Figure 4a reports the sample size needed to investigate this effect, with an alpha of 0.05 and power of 0.9 (given the high uncertainty of the field the safest approach would be to plan a high power). The randomised controlled trials are analysed in order of publication time, with the effect size of the (theoretical) living meta-analysis being updated every time a new trial is published. As shown in figure 4, all updates of the living meta-analysis are reporting non-significant results that never cross the monitoring boundaries (lack of evidence), and that the APIS needed (given the minimal clinically relevant RRR, alpha and power) was not reached. Had a living meta-analysis been planned and performed, it would have shown that all published CBT trials were underpowered for testing the prevention of psychosis at 6-month follow-up under the above parameters (i.e. were false negatives). Figure 4b investigates the preventive effect of CBT versus needs-based interventions in CHR-P individuals at 12 months. With the second trial there are significant results but a spurious effect because the curve does not cross the monitoring boundary (false positive result). The fourth and fifth trials produce significant results that cross the monitoring boundaries before the information sample is reached (true positive results). However, with the addition of the sixth and seventh trials the required APIS is almost met, the monitoring threshold is not crossed, and the result becomes non-significant (this could be a true negative result). Importantly, as noted above, these estimates relate to relatively large effect sizes (RR=0.5). Current interventional CHR-P literature to date has not been powered to detect smaller effect sizes.

DISCUSSION
This is the first umbrella review of the evidence for the effect of preventive interventions on several clinical outcomes in CHR-P individuals. Seven meta-analyses that were investigating preventive treatments in CHR-P individuals were included. None of them produced pooled effect sizes across psychological, pharmacological or other types of interventions. The outcomes analysed encompassed: risk of psychosis onset from a CHR-P state, acceptability of treatments, severity of attenuated positive and negative psychotic symptoms, depression, symptom-related distress, social functioning, general functioning, and quality of life. There was no evidence to favour any of the current preventive interventions over each—and any—other to improve any of these clinical outcomes. These meta-analyses were based on 20 randomised controlled trials. The vast majority of them (70%) defined the prevention of psychosis onset as their primary outcome (first-order issue) of interest. Several new trials are ongoing in this population, reflecting the high interest in this field. Simulation of a living meta-analysis did not change the interpretation of the current evidence.

The following sections interpret these findings in the context of the overall evidence for interventions in CHR-P individuals, the use of pooled vs treatment-specific effect sizes, the definition of primary and secondary outcomes in the published trials and future recommendations to advance clinical research in this field.

**Evidence-based efficacy of preventive treatments for CHR-P individuals**

As noted in the introduction, it has been argued that the CHR-P paradigm has the capacity to (i) identify people at incipient risk for psychosis, (ii) improve their levels of distress and functioning, and (iii) reduce their risk of progression to sustained psychotic disorder[32]. It is true that the CHR-P paradigm has allowed the first prospective identification of individuals at risk of developing psychosis in clinical psychiatry. The achievements of the CHR-P paradigm have been extensively celebrated over the past years[3]. At the same time, some important challenges have emerged. As illustrated by the trough of disillusionment phase in the Hype Cycle in Figure 1, the actual ability of specialised CHR-P clinics[4] to detect individuals at risk for psychosis is limited. An original study has estimated that only around 5% of individuals who will later develop a first episode of psychosis in secondary mental health care were detected by the local services at the time of their CHR-P stage[15; 91]. Front-line youth mental health services, as opposed to specialised CHR-P clinics
such as the OASIS, can be expected to detect more at-risk individuals. However, the limited evidence indicates that even youth mental health services can detect only 12% of first episode cases at the time of their CHR-P stage[92]. It is therefore clear that we need to improve our ability to detect CHR-P individuals in secondary mental health care, primary care and in the community. Some experimental approaches have been validated, which include automatic risk calculators for use in mental health trusts[57; 91; 93] or in the community[94].

Conversely, there is no evidence that current treatments are effective in improving the level of distress and functioning of CHR-P individuals. The most updated meta-analytical evidence summarised in Table 2 shows no evidence to favour any preventive treatments over each (and any) other for improving any of the clinical outcomes that are relevant for CHR-P individuals: risk of developing a first episode of psychosis, acceptability of treatments, severity of attenuated positive or negative psychotic symptoms, depression, symptom-related distress, level of social functioning, level of general functioning and quality of life. Although CHR-P patients may improve in their symptoms pre-post treatment, there are no substantial differences across interventions (which also includes control conditions). Importantly, although the meta-analyses presented in Table 2 were conducted by different and independent research teams, their results converge. For example, two independent meta-analyses both concluded that there is no evidence to favour specific treatments over each other for improving attenuated positive psychotic symptoms in CHR-P individuals[58; 63]. Unfortunately, there is no robust evidence that current treatments can improve functional levels in these patients. The most recent meta-analysis to have explored this outcome concluded that Cognitive Behavioural Therapy did not significantly improve social functioning at 6 months, 12 months or 18 months; Omega-3 did not significantly improve social functioning at 6 months or 12 months and Cognitive Remediation did not significantly improve social functioning at 2- to 3-month follow up[65]. Lack of evidence for a symptomatic effect of preventive treatments in CHR-P individuals is also in line with our limited understanding of their mechanism of action at a neurobiological or psychological level[95].

**Apples, oranges and the production of clinically meaningless pooled effect-sizes**
A second issue, as detailed in the introduction[32], is the possibility that if the two recent network meta-analyses[30; 63] had pooled together psychosocial and pharmacological trials, statistically significant findings could have emerged[32]. On a statistical and conceptual level, if the randomised controlled trials are all pooled together, there could be no “network” meta-analysis comparing specific nodes against each other but just a simple meta-analysis of the average efficacy of all treatments. On a pragmatic level, it is indeed true that these two meta-analyses[30; 63] did not even estimate a pooled effect size for all types of treatments, because this has little clinical interest and, as such, it was deliberately not planned in the a priori protocol of these studies[96]. These meta-analyses were conducted to estimate the level of evidence with respect to specific clinical questions. Since preventive treatments for CHR-P individuals are highly heterogeneous, pooled effect sizes are clinically meaningless and cannot be used to inform treatment guidelines. How would a clinician, who needs to decide on the best treatment to offer to a CHR-P patient, fare when interpreting a pooled effect size which has been estimated across pharmacological, psychological and dietary interventions vs heterogeneous control conditions? Such an effect size would, at best, suggest that “any type of experimental treatment” is better/worse/similar than “any type of control condition”. This is some information (i.e. the intervention seems on average to work), but it is clearly useless from a clinical standpoint. Treatment guidelines need to be as specific as possible because clinicians, healthcare providers and patients rely on them for clear direction and education on the best ways of adhering to them. Three previous aggregate pairwise meta-analyses were cited by the authors as support for pooling effect sizes across pharmacological and psychological categories of treatments[32]. The first meta-analysis cited did not estimate any pooled effect sizes at all[26]. Rather, this meta-analysis performed separate pairwise comparisons of specific preventive interventions, such as cognitive behavioural therapy vs supportive counselling[26]. The second meta-analysis did compute significant pooled effect sizes and concluded that “receiving any of the focused treatment[53]” was associated with a reduced risk of psychosis onset. This clearly demonstrates the criticalities above: what exactly is a “focused treatment” and how should it be implemented in clinical routine? Because the authors could not answer such a question, they stated that their results did “not allow recommendation for any specific treatment”[53]. This further demonstrates that pooled effect sizes in the CHR-P field are clinically meaningless and cannot inform
treatment recommendations. It is, therefore, important to examine the efficacy of specific treatment effects in CHR-P individuals. Accordingly, the rationale for conducting the first randomised controlled trial in the CHR-P population was based on the need to examine more “specific interventions” – page 922 in [19]. The third meta-analysis which was cited did compute a pooled effect size across cognitive behavioural therapy plus risperidone, olanzapine, omega-3, integrated psychotherapy (which included group skills training, cognitive remediation and multifamily psychoeducation in addition to cognitive behavioural therapy), and cognitive behavioural therapy compared to treatment as usual [52]. In this case, the overall effect size was reported in figure 2 on page 60 [52] and it was inaccurately termed, in the figure itself, as “cognitive behavioural therapy” vs “treatment as usual”, ignoring the effect of risperidone, olanzapine and dietary interventions [52]. Furthermore, when this meta-analysis stratified the effect size by type of interventions, it concluded that “the subset of CBT-based interventions is associated with a pooled RR of 0.52 (95% CI = 0–79)”, and reported 95% confidence intervals which included the null hypothesis (RR=1) [52]. The lack of statistical significance could be due to the lower statistical power because of the stratification, yet this finding does not support a robust preventive effect of CBT-based interventions. This demonstrates that pooled effect sizes are not only of difficult clinical interpretability but also at very high risk of reporting biases.

Interventions for CHR-P are intrinsically highly heterogeneous because they include different therapeutic components. This is due to the fact that when interventions were originally introduced in the first randomised controlled trial, the strategy adopted was to “include the best-bet specific therapies in a single enhanced intervention package to determine whether it was possible to delay the onset of psychosis” [19]. The subsequent randomised controlled trials have followed this approach by testing different packages, each of which was characterised by specific therapeutic components. This issue is evident, for example, in the case of different types of psychological therapies, which have been defined as “black boxes” [97]. The additional and substantial problem with using pooled effect sizes in the CHR-P field is that the control group, traditionally termed as “treatment as usual” [52], is per se poorly standardised and largely dependent on local service configurations and the availability of specific resources or competences [61]. For example, treatment as usual may encompass supportive psychotherapy primarily focusing on pertinent issues such as
social relationships and vocational or family problems, case management, providing psychosocial assistance with accommodation, education or employment, brief family psychoeducation and support, medications other than antipsychotics or clinical monitoring and crisis management[26]. One way to overcome these issues is to employ network meta-analyses. Network meta-analyses offer additional benefits over standard pairwise analyses in that the comparative efficacy of specific interventions can be estimated and ranked, even when two treatments have never been compared directly head-to-head[98]. Furthermore, network meta-analyses can improve the precision of estimates by allowing integration of both direct and indirect treatment effect estimates[99]. The World Health Organization recommends network meta-analyses over pairwise meta-analyses as a basis for informing clinical guidelines[100]. Therefore, network meta-analyses should be considered the highest level of evidence in CHR-P treatment guidelines[101]. Because of these reasons, over recent years independent research teams have used network meta-analytical approaches to investigate the specific efficacy of preventive interventions for CHR-P individuals. Importantly, the current umbrella review demonstrates that none of them has produced pooled effect sizes[58; 60; 62; 65]. This further provides independent validation that pooled effect sizes across different types of treatments should not be estimated when conducting treatment meta-analyses in the CHR-P field.

The opportunistic (in)significance of preventing the onset of psychosis
The third issue raised by some authors was that recent meta-analyses have focused on “second-order issues”, namely transition to psychosis in CHR-P individuals[32]. The relegation of the prevention of psychosis to a second-order clinical outcome conflicts with the previous two decades of experimental research in this field. The extent to which randomised controlled trials in CHR-P individuals have prioritised prevention of psychosis, compared to other outcomes, is reported in Table 2. All twenty randomised controlled trials but four, corresponding to 70% of published trials, have focused on preventing the first onset of psychosis in CHR-P individuals as their primary outcome. Even the trials that did not primarily focus on preventing transition to psychosis still recommended investigation of whether the specific interventions could affect conversion rates[67] and modify the course of disease[66].
Until now, there has been a converging consensus in the CHR-P field that prevention of psychosis was the foremost outcome and the ultimate aim of the entire paradigm. Authors of randomised controlled trials in CHR-P individuals have declared that the goals of early detection are: 1) postponement or prevention of the transition to frank psychosis, 2) reduction of the duration of untreated psychosis to a minimum in patients who develop florid psychosis, and 3) prevention of delay to accessing mental health services[102]. The authors also explicitly acknowledged that prevention of psychosis is the main goal[102]. In line with this, outcomes other than the onset of psychosis have been poorly operationalised in the CHR-P field. For example, there is no clear definition for symptomatic remission or good outcomes in CHR-P patients. The importance of preventing the onset of psychosis in CHR-P individuals has been further endorsed worldwide and supported by consensus papers[3]. Notably, the first randomised controlled trial in these patients aimed at determining whether it was possible to delay the onset of psychosis[19]. The rationale given was overall pointing to the fact that this would be the most important way to alter the course of the disorder and eventually improve the lives of many patients. Such accumulating evidence clearly indicates that the prevention of psychosis has been “the” first-order issue in the CHR-P and not at all a second-order issue, as argued[32]. It does not seem justified to downgrade the relevance of preventing psychosis in CHR-P individuals just because recent meta-analyses have not found robust evidence to favour specific preventive interventions (including also control conditions such as needs-based interventions). It has been argued that recent meta-analyses are “pessimistic” and have “distracted from the key message” of the research literature, which is “in need of positive findings pointing to better outcomes”[32]. Evidence synthesis is conducted to test the robustness of findings on a determinate topic, and not to meet the needs of scientists and researchers. Reporting negative as well as positive findings is equally important. By systematically identifying and addressing gaps in knowledge, “negative” evidence syntheses such as these may, in fact, help to advance the field. For example, in taking a closer look, recent meta-analyses have concluded that owing to large confidence intervals, the actual efficacy of treatments for preventing psychosis is mostly indeterminate[30]. Some effective signal in treatments’ efficacy may have been missed because of the large clinical heterogeneity[61] of the population being investigated. In this sense, these results are not at all pessimistic, but they rather call for a new generation of experimental research in this field (see below). Conversely,
the claim that prevention of psychosis is a second-order issue could discourage future trials from exploring this relevant outcome and the current uncertainty in the meta-analytical estimates may remain unresolved. On a more conceptual level, there is some consensus that psychosis onset as defined categorically is an arbitrary concept[103; 104; 105] and that it should be better refined or complemented by other relevant outcomes, such as severity of attenuated symptoms, disability and functional outcomes. Yet, prevention of psychosis should remain the cornerstone and the most important outcome for the CHR-P field, complemented by other outcomes such as severity of symptoms, functional improvement, recovery and quality of life. Downgrading—or at worst, dropping—such a primary outcome would be an indirect demonstration that current CHR-P research has approached the lowest depths of the through of disillusionment, with complete failure with respect to its ability to prevent psychosis. It would also prove that preventive psychiatry is different from other branches of medicine. For example, lack of effective treatments to prevent dementia[106] from a mild cognitive impairment stage has not triggered neurologists to claim that prevention of dementia itself should be downgraded to a second-order issue (instead it has become a priority for public health)[107].

Towards the slope of enlightenment and the next generation of experimental therapeutics

As noted above, some authors have used recent findings to claim that the CHR-P paradigm should be completely abandoned. Such a claim seems too intransigent and not evidence-based. For example, by using the same criteria we should claim that integrated clinical services for first episode of psychosis should be dismantled, given that there is no evidence that they can reduce the duration of untreated psychosis[108] or prevent relapse[1]. The absence of evidence is not evidence of absence[109]. Furthermore, dismissing two decades of CHR-P research is not in line with the progressive nature of medical knowledge. As clarified in Figure 1, the trough of disillusionment often precedes the slope of enlightenment and the plateau of stable knowledge. The uncertain stage of knowledge which is typically associated with the trough of disillusionment is not specific to the CHR-P state; rather, it has also been observed in other branches of clinical medicine, such as cancer prevention[110]. It is thus important to use our understanding of current limitations to advance knowledge, rather than hinder it.
The essential role of risk enrichment during recruitment

The psychometric instruments traditionally used to ascertain the presence of a CHR-P state have been validated worldwide and, overall, have demonstrated excellent prognostic accuracy (AUC at 3 years: 0.9), comparable to that of other prognostic instruments employed in organic medicine[7]. Yet, such an excellent prognostic accuracy is mostly due to the excellent sensitivity (96%) of CHR-P instruments to detect a state of risk for psychosis. Therefore, they are able to detect nearly all individuals who would develop psychosis, and consequently, nearly all individuals testing negative should be individuals who would not develop psychosis. In other words, a negative CHR-P assessment is associated with a very small probability of developing psychosis (1.56% at 3-year, negative likelihood ratio of 0.09)[7; 111]. Conversely, the CHR-P instruments have a poor specificity (47%) and thus half of the individuals who would not develop psychosis might be false positives. Therefore, among individuals testing positive, most would not develop psychosis, or in other words, a positive CHR-P is still associated with a relatively small probability of developing psychosis (26% at 3-year, positive likelihood ratio of 1.82)[7; 111]. Therefore, there is only a limited predictive gain in testing positive at a CHR-P assessment. As a consequence, CHR-P instruments should be used in samples that have already been enriched in their initial risk of psychosis and not as screening methods in the general population (see below)[7; 112]. Otherwise, the global number of false positives would be so high, that most individuals with positive testing would be false positives[112]. When individuals undergoing a CHR-P assessment are recruited from mental health services, they accumulate several risk factors for the disorder[6] which increase their level of risk to 15% at 3 years, compared to the 0.43% 3-year risk in the local age-matched general population[9; 113]. This level of risk is also termed “pretest risk” because it is ascertained in the whole group of people undergoing a CHR-P assessment before the results of the assessment itself are known[111]. The pretest risk in individuals recruited through mental health services (i.e. measured in naturalistic studies, excluding randomised controlled trials) is 15% at 3 years, worldwide[5]. However, such an estimate is highly variable, with study estimates ranging from 3% to 49%, because it is based on unstandardised idiosyncratic recruitment strategies[5].
fold increase from the pretest of 15%) and those not meeting CHR-P criteria will have a 1.56% risk of developing psychosis at 3 years (10-fold decrease from the pretest risk of 15%). As indicated in Table 4, assuming an alpha of 0.05, power of 90% (a higher power of 90% is recommended given the high uncertainty of this field; see eTable 1 for sample size estimations using 80% power), 2-sided test, it is possible to estimate the sample size which is required to test a new experimental treatment to prevent psychosis against treatment as usual (e.g. needs-based intervention). If the experimental treatment is able to halve the risk of psychosis (risk ratio=0.5), 2368 CHR-P individuals (conservatively using the 3% lower bound of the study estimates to avoid additional underpowered negative trials[31]) are required to complete the trial. After considering some attrition due to lost to follow-up (e.g. 20%), the final sample size would be of 2842 CHR-P individuals. The problem is that this sample size is based on the pretest estimate from naturalistic studies that more likely represent the whole population seeking help at specialised CHR-P clinics. When CHR-P individuals detected by these specialised clinics are recruited into randomised controlled trials, it is likely that additional sampling biases would apply, further reducing the risk enrichment. For example, the push towards recruiting sufficient numbers of trial participants within a fixed period of time may lead to more intensive outreach campaign in the local community, which are well known to dilute the pretest risk[114]. In other words, using unstructured recruitment strategies it may not be sufficient to recruit 2842 CHR-P in new trials to be sufficiently powered to test preventive effects of magnitude 0.5. Furthermore, such a study would not be powered to test smaller effect sizes that are likely in the case of specific treatment components. If the efficacy of the preventive is lower, for example, if the experimental treatment is able to reduce the risk of psychosis onset by only 40% (risk ratio =0.6) or 30% (risk ratio=0.7), 3942 and 7436 CHR-P patients are needed to complete the trial (ignoring attrition), respectively. If individuals undergoing a CHR-P assessment are mostly recruited from the community, they will have accumulated less or no risk factors for psychosis[61] and their pretest risk would be 0.43% at 3 years. Following the estimates reported in Table 4, the sample size required for a similar randomised controlled trial would exceed 17908 CHR-P individuals (ignoring attrition). In fact, CHR-P instruments do not work well when they are applied outside clinical samples that have already undergone some pretest risk enrichment[112; 114]. As noted above, the way individuals are recruited (for undergoing a CHR-P assessment) drives the level of risk
and ultimately, impacts the statistical power of the trial. For example, the Neurapro trial has observed a risk of psychosis onset of about 14\% at 3.4 years[29] in CHR-P individuals who had received the control condition, which suggests a pretest risk of 8.2\% (assuming that the control condition had little effect on the risk to psychosis). Under those circumstances, 796 CHR-P individuals would be needed to detect a 50\% decrease in risk in the experimental condition, which exceeded the sample size employed by the trial. In the case of some effect of the control condition on the risk of transitioning to psychosis (as speculated by the authors[28]), the estimated sample size would then be lower.

It is therefore possible that the low level of risk for psychosis decreased the statistical power of the trial to detect small signal effects associated with the experimental treatments. Lack of statistical power because of a poor level of psychosis risk may actually be one of the causes of the negative randomised controlled trials in this population[31] and of the associated large 95\% confidence intervals that have been observed in the last network meta-analysis[30]. The main problem is that recruitment strategies in this field are idiosyncratic and poorly standardised and as such, it is not possible to control the level of pretest risk enrichment. This is particularly concerning in the case of recruitment into trials, which, as noted above, introduces additional selection biases and may further dilute the risk enrichment. For example, likely because of more intense outreach campaigns in the community, the actual risk of psychosis in CHR-P samples has been declining from 29\% (2012[115]) to 20\% (2016[12]) worldwide. Interestingly, there are exceptions to this phenomenon, such as the Outreach and Support in South London (OASIS) CHR-P service[4], where transition risk has not declined over time. This is again due to the fact that recruitment strategies have, overall, maintained a stable pretest risk enrichment[116]. Ultimately, innovative strategies are needed to ensure that a sufficient level of risk enrichment is obtained to allow adequate statistical power in CHR-P recruited into future trials. A possible solution could be to apply pretest risk stratification algorithms that have been developed and validated for this population using machine learning methods (Figure 5)[113]. Machine learning methods generally represent innovative techniques that are expected to deliver more robust prognostic models[117]; their real-world impact in the field, however, awaits further confirmation[118].
For example, pretest risk estimation algorithms that are based on the source of referral to CHR-P clinics and ethnicity can be used to stratify individuals into four classes of risk enrichment: low risk (1% risk at 3 years, ≈21% of the CHR-P population), moderately low risk (10% at 3 years, ≈53% of the CHR-P population), moderately high (18% at 3 years, ≈21% of the CHR-P population), and high (46% at 3 years, ≈5% of the CHR-P population, Figure 5). If this simple tool is applied to individuals recruited for a CHR-P assessment it may be possible to control risk enrichment and avoid its high variability. Furthermore, stratification may inform the subsequent phases. For example, those who are at low risk are to be screened out from trial eligibility, a pretest risk enrichment of at least 10%—or more—would be ensured across all participating sites (Figure 5). Under those circumstances, a total sample of 648 CHR-P individuals (ignoring attrition) would guarantee sufficient statistical power (90%) to test treatment effects that can halve the risk of developing psychosis. Alternatively, those with a moderately low pretest risk can be subjected to further testing to refine the prognostic accuracy[119]. Controlling pretest risk enrichment through the recruitment phase is expected to mitigate the existing challenges and facilitate the slope of enlightenment phase (Figure 1).

**Stratification and precision medicine**

As noted above, the lack of evidence to favour specific preventive treatments over any others should be the basis to promote further research in this field, rather than to abandon it. For example, it is likely that a one-size-fits-all treatment approach in CHR-P populations is not effective and that some treatments may work for specific subgroups of patients. There are several lines of evidence to support this. First, some meta-analyses included in the current umbrella review report large 95% confidence intervals, which indicate high heterogeneity across treatments. Some of them may actually be effective. For example, the network meta-analysis exploring the effect of treatments on the prevention of psychosis has suggested that the integrated psychological intervention developed by Bechdolf et al[22] has the largest effect size for reducing the risk of psychosis onset compared to needs-based interventions (Odds Ratio=0.04)[30]. Although the 95% confidence intervals included the null hypothesis, this treatment could be the focus of the next generation of therapeutic trials. An important note is that current meta-analytical evidence did not favour any treatment
compared to each other (including control conditions such as needs-based-intervention). As such, it is possible that needs-based-intervention per se was associated with improved outcomes, diluting the potential comparative efficacy of other interventions. Another example comes from the failure of some randomised controlled trials, such as the NEURAPRO (omega-3) when tested in an unstratified CHR-P sample[28]. This has naturally led to the suggestion that omega-3 might be more efficacious in those individuals who specifically have low levels of membrane fatty acids at baseline[120]. This makes sense from a pathophysiological perspective—psychosis is a heterogeneous disorder with likely many different neurobiological ‘paths’ and risk factors[61], which may perturb an individual’s neural circuitry in various (innumerable) ways and which ultimately presents as psychosis[95].

Table 5 provides a clinical interpretation of the evidence for selecting preventive interventions for CHR-P individuals.

Second, there is converging evidence indicating that the CHR-P group is clinically heterogeneous. For example, there is high heterogeneity in the level of risk for psychosis across the three CHR-P subgroups of the attenuated psychotic symptoms, brief and limited intermittent psychotic symptoms and genetic risk and deterioration syndrome[12; 104; 105]. In particular, those meeting the brief and limited intermittent psychotic symptoms criteria have a very high risk of developing persistent psychotic disorders. Recent studies also demonstrate that these patients are those in highest clinical need but receive inadequate care[121; 122]. Such a heterogeneity calls for a revision of the CHR-P paradigm which should include clinical stratification across these three subgroups[10; 17]. Ongoing international consortia such as PSYSCAN, PRONIA and NAPLS have already started delivering precision medicine tools for stratifying CHR-P individuals and may permit an individualised prediction of their outcomes[57; 123; 124]. These consortia are adopting a multimodal prediction approach which is spanning clinical, neurocognitive, neurobiological as well as genetic predictors[125] to improve the individualised prognosis of outcomes in this population.

**Individual participant data living meta-analyses**

Another precision medicine approach that is expected to advance knowledge is to conduct further evidence synthesis studies such as individual-participant data level
network meta-analyses. The use of individual data as opposed to aggregate-level meta-analysis would also allow analysing the effect of different individual factors (including the CHR-P subgroups) and development of evidence-based prognostic algorithms to forecast the likelihood of treatment response at the individual subject level based on these factors. It is expected that these individual-participant data meta-analyses will identify specific subgroups of CHR-P individuals for whom current treatments (i.e. the integrated psychological intervention) may already be effective. A relevant issue (highlighted in Table 5), is that the future publication of a single study with robust evidence of efficacy may significantly change the level of evidence for preventive interventions in psychosis. As indicated in table 3 and in the section below, novel compounds for this patient population are under investigation, and it is thus expected that new results will be released over the next few years. This would be the ideal context to conduct living meta-analyses. To illustrate their methodological approach and the potential impact in the field we have simulated a living meta-analysis. The simulation showed no robust evidence to favour current CBT treatments for the prevention of psychosis. Prospectively planned living systematic reviews and meta-analyses are thus expected in this field. The feasibility and sustainability of living systematic reviews can be supported by the synergic interaction of human effort and machine automation[126]. A recent empirical study has demonstrated that prospectively planned living network meta-analyses produced strong evidence against the null hypothesis more often—and earlier—than conventional pairwise meta-analyses[39].

Innovative youth mental health services and advanced trial designs
Additional important factors that may support the slope of enlightenment are the development of youth mental health services and innovative trial designs. The high prevalence of mental health problems among young people, their negative impact on outcomes, as well as their significant financial and societal cost, emphasises the need to improve broad mental health care in children and young people[127]. The ‘No Health Without Mental Health’ report by the UK Government recognised that only a life-course approach would allow fulfilment of such an objective, and highlighted the importance of the early years[128]. Similarly, the ‘Future in Mind’ report by NHS England highlights the pressing need (by 2020) for a holistic approach, including better access and support for front-line staff, and the adoption of innovative integrated youth
mental health approaches which depart from the current tier system which is split between Child and Adolescent Mental Health Services and Adult Mental Health Services[129]. The OASIS model presented above already provides integrated care across child and adolescent and adult mental health services for CHR-P individuals aged 14-35. Notably, the CHR-P services platform has been recently implemented nationwide in some countries, demonstrating scalable impact for taking care of children and young adults. Therefore, it may be possible to leverage these CHR-P templates in order to refine the next generation of youth-friendly mental health services which target the needs of adolescents and young adults experiencing early stages of other mental disorders, such as depression[130] and bipolar disorders[131; 132]. Some integrated models of care have already capitalised on the CHR-P platform to broaden their horizons and target the wider mental health of children and young adults.

In 2006, following a campaign that was led by leaders in mental health, the early intervention model for psychosis was then expanded to other diagnoses (e.g., mood, personality, eating and substance use disorders) through the creation of Headspace in Australia[133]. Headspace is a government-funded initiative that provides youth-friendly, stigma-free early intervention services in a ‘one-stop shop’ location to 12–25-year olds with emerging mental health and substance use disorders[133]. Within this context, mental healthcare is multidisciplinary, integrated, delivered in a single setting and is centred on the needs of young people along with their families[134]. The youth mental health reform achieved in Australia has permeated to other regions of the world, with the UK, Ireland, Canada, USA, Europe and Asia adopting similar, culturally appropriate models[134; 135]. Youth mental health services may offer some additional clinical research advantages because they fully embed the clinical staging model for the emergence of several mental disorders[1; 136]. In turn, this would facilitate stepped clinical care. For example, by adopting subsequent prognostic assessments[8] in young people at risk for emerging mental disorders, it may be possible to refine the prediction of their longitudinal outcomes whilst optimising the logistic resources and translational impact[119]. The stepped prognosis can be associated with stepped care, which targets the level of need and risk presented by each person. Adaptive clinical trials represent an innovative and flexible method, allowing pre-specified modifications to the design or statistical procedures of an on-going trial depending on the data generated from it. Such adaptive design trials can boost clinical research by cutting
costs and time, and more importantly, by adapting to the clinical stages of the stepped care model implemented in youth mental health services. The first adaptive design clinical trial, the Staged Treatment in Early Psychosis Study (STEP)[137] is ongoing, and more are expected to follow. However, although holistic packages of care that are embedded in new mental health services promise to deliver substantial therapeutic advancements, the effectiveness of their specific treatment components still need to be carefully assessed and evaluated.

New experimental therapeutics
As indicated in our latest network meta-analyses[30; 63] and in Table 3, innovative experimental therapeutics in CHR-P individuals are under development. These compounds, if used along with the above strategies are expected to advance knowledge in this field.

In summary, the slope of enlightenment and future advancements of knowledge in this field could be facilitated by controlling pretest risk enrichment in samples undergoing CHR-P assessment, stratifying CHR-P samples to allow precision medicine approaches, conducting Individual Participant Data and living meta-analyses, adopting innovative youth mental health services and advanced trial designs and testing novel experimental therapeutics.

CONCLUSIONS
This umbrella review found no convincing evidence for superior efficacy of any current preventive treatment, compared to each—and every—other, on the risk of psychosis onset, acceptability of treatments, severity of attenuated positive and negative psychotic symptoms, depression, symptom-related distress, social functioning, general functioning, or quality of life in CHR-P individuals. Current interventional literature in CHR-P population has been only powered to detect large effect sizes for preventive treatments. Prevention of psychosis from a CHR-P state has been and should remain the (first order) primary outcome of interventional research, refined and complemented by other clinically meaningful outcomes. The stagnation of knowledge in the field should promote innovative and collaborative research efforts, rather than give us cause to abandon it. Advancements will most likely be associated with the ability to deconstruct the high heterogeneity of CHR-P populations. This would require
the estimation of treatment-specific effect sizes through individual participant data meta-analyses, controlling risk enrichment during recruitment and embedding stratification and precision medicine methods within new youth mental health services that can accommodate staged care, stepped prognosis and advanced trial designs. Several experimental therapeutics studies in CHR-P individuals are also ongoing. The strategies which could facilitate reaching the plateau of knowledge in CHR-P field are summarised in Gartner Hype Cycle (Figure 1). We hope that these innovations will ultimately pave the way toward a future plateau of knowledge and refined preventive treatments that could result in tangible benefits for clinicians, researchers and, more importantly, for patients and their families.
Figure 1. The hype cycle of preventive treatments for psychosis

Innovation trigger

Peak of inflated expectations
- NICE CG178 guidelines, 2014
- EPA guidelines, 2015

IPI trial
- Bechdolf et al 2012

Omega 3 trial
- Amminger et al 2010

Olanzapine trial
- McGlashan et al 2004

CBT trial
- Morrison et al 2004

Risperidone+CBT trial
- McGorry et al 2002

First CHR-P clinical service
- Yung et al 1995

Largest CBT trial negative
- Morrison et al 2012

Largest Omega 3 trial negative
- McGorry et al 2017

Few at-risk people detected by CHR-P clinics
- Fusar-Poli 2017

Trough of disillusionment

No meta-analytical evidence impacting:
- Risk of psychosis onset (Davies 2018)
- Symptoms/functioning (Devoe 2018)

Slope of enlightenment

Plateau of knowledge

Innovative trial designs
- New experimental treatments
- Youth mental health services
- IPD & Living meta-analyses
- Stratification & Precision Medicine
- Recruitment & risk enrichment

Visiblity

Time (years)
Figure 2. Living meta-analyses. Current (inner circle) and emerging (outer circle) evidence-based health knowledge ecosystems. The current health knowledge ecosystem is characterised by inefficiencies that hamper the flow of knowledge from health practice through primary research, evidence synthesis and guidelines, and finally back to impacts on health practice. The emerging health knowledge ecosystem is characterised by a continuous flow of knowledge between living components, including the growing importance of learning health care systems (a dynamic system which is continuously learning from new data), which together with traditional primary research will populate common data repositories. Living evidence services derived from these repositories, supporting living guidance and decision support systems will close a “living” health knowledge loop. Adapted from[41].
Figure 3. Study identification and selection (PRISMA flowchart)

Records identified through database searching (n = 2328)

Meta-analyses (n = 327)

Records screened on the basis of title and abstract reading (n = 327)

Records excluded (n = 313):
- not focusing on CHR patients
- not focusing on interventions

Full-text meta-analyses assessed for eligibility (n = 14)

7 full-text articles excluded:
- No quantitative data reported (n=1)
- Overlapping old meta-analyses (n=6)

Meta-analyses included in the umbrella review (n=7)
Table 1. Efficacy of treatments for CHR-P individuals. Overview of the most recent meta-analyses per clinical outcome (up to January 11th, 2019).

| Outcome                                                        | Author                        | year  | Type of evidence                        | N of studies (max n of CHR individuals) | AMSTAR rating | Finding                                                                 |
|---------------------------------------------------------------|-------------------------------|-------|-----------------------------------------|------------------------------------------|---------------|-------------------------------------------------------------------------|
| Transition to psychosis                                       | Davies et al[30]              | 2018  | Aggregate Network Meta-Analysis (RCTs)  | 16 (2035)                                | 10/11         | Lack of evidence to favour specific treatments                          |
| Acceptability                                                 | Davies et al[30]              | 2018  | Aggregate Network Meta-Analysis (RCTs)  | 14 (1848)                                | 10/11         | Lack of evidence to favour specific treatments                          |
| Severity of attenuated positive psychotic symptoms            | Davies et al[63]              | 2018  | Aggregate Network Meta-Analysis (RCTs)  | 14 (1707)                                | 10/11         | Lack of evidence to favour specific treatments                          |
| Severity of attenuated negative psychotic symptoms            | Devoe et al[58]               | 2018  | Aggregate Network Meta-Analysis (RCTs)  | 12 (1457)<sup>b</sup>                    | 10/11         | Lack of evidence to favour specific treatments                          |
| Depression                                                    | Stafford et al[26; 59]         | 2013  | Aggregate Pairwise Meta-Analysis (RCTs) | 5 (714)                                  | 9/11          | No significant treatment effects at any time point                      |
| Symptom-related distress                                      | Hutton et al[64]              | 2014  | Aggregate Pairwise Meta-Analysis (RCTs) | Unclear                                  | 9/11          | No significant treatment effects                                         |
| Social functioning                                            | Devoe et al[65]               | 2018  | Aggregate Pairwise Meta-Analysis (RCTs) | 9 (1040)                                 | 10/11         | No treatment significantly improved social functioning                  |
| Functioning                                                   | Schmidt et al[25]             | 2015  | Aggregate Pairwise Meta-Analysis (RCTs) | 9 (869)                                  | 8/11          | No significant treatment effects                                         |
| Quality of life                                                | Hutton et al[64]              | 2014  | Aggregate Pairwise Meta-Analysis (RCTs) | Unclear                                  | 9/11          | No significant treatment effects                                         |

<sup>a</sup> sample sizes are based on the total sample sizes reported in the meta-analysis minus the sample size of any studies that were not included in their actual meta-analytic computations.

<sup>b</sup> sample size of Ising et al (2016) and the non-randomised arm of McGorry et al 2013 (N=78) not included.

<sup>c</sup> sample size computed by summing study sample sizes from Table 1. The non-randomised arm of McGorry et al 2013 (N=78) was not included.
| Author                      | Primary outcome          | Rationale supporting the primary outcome                                                                                                                                                                                                 |
|-----------------------------|--------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Addington et al 2011[138]   | Prevention psychosis     | Psychological interventions might be expected to be promising in the pre-psychotic period when the symptoms are less severe and also less specific                                                                                                           |
| Amminger et al 2010[23]     | Prevention psychosis     | Intervention in at-risk individuals holds the promise of even better outcomes, with the potential to prevent full blown psychotic disorders.                                                                                                                                               |
| Bechdolf et al 2012[22]     | Prevention psychosis     | Prevention efforts in individuals at imminent risk of schizophrenia can reduce or prevent the devastating effects of the disorder                                                                                                                                                  |
| Bechdolf et al 2016[139]    | Prevention psychosis     | Effective interventions for CHR-P individuals are needed in order to reduce or prevent the devastating effects of the disorder                                                                                                                                         |
| Cadenhead et al 2017[140]   | Prevention psychosis     | Replication study testing the efficacy of dietary interventions as defined by Amminger et al[23]                                                                                                                                                                                               |
| Choi et al 2017[71]         | Social functioning       | Providing cognitive remediation during a putative prodromal stage may improve social functioning and have some value in reducing the risk of psychosis onset                                                                                                                                 |
| Kantrowitz et al 2015[66]   | Reduction of latent psychotic symptoms | Negative symptoms and cognitive deficits frequently persist and contribute substantially to impaired functional outcome.                                                                                                                   |
| Loewy et al 2016[70]        | Cognitive functioning    | The variability in outcomes for CHR-P patients requires treatments that offer the prospect of high benefit and low risk                                                                                                                                           |
| McGlashan et al 2006[20]    | Prevention psychosis     | The chronicity of schizophrenia determines the primary rationale for studies of early intervention for this disorder                                                                                                                                                        |
| McGorry et al 2002[19]      | Prevention psychosis     | Progression to psychosis is neither inevitable nor predetermined and it may be possible to delay the onset of psychosis                                                                                                                                                         |
| McGorry et al 2017[28; 29]  | Prevention psychosis     | Treatment strategies should relieve distress, improve functioning, and reduce the risk for progression to a psychotic illness                                                                                                                                                    |
| Miklowitz et al 2014[67]    | Reduction of latent psychotic symptoms | Intervention during the high-risk period may reduce subthreshold psychotic symptoms, enhance social and role functioning, and, over the long term, prevent or delay conversion to episodes of psychosis |
| Morrison et al 2004[21]     | Prevention psychosis     | Specific pharmacotherapy and psychotherapy may be associated with a reduction in progression to psychosis in CHR-P people                                                                                                                                                   |
| Morrison et al 2012[27]     | Prevention psychosis     | Effective interventions to prevent or delay this transition are needed because of the significant personal, social, and financial costs associated with the development of psychosis                                                                                           |
| Piskulic et al 2015[69]     | Cognitive functioning    | Given that deficits in cognition are related to poor functional outcome in CHR-P, cognition is a good treatment target                                                                                                                                                      |
| Ruhrmann et al 2007[68]     | Reduction of latent psychotic symptoms | Attenuated psychotic symptoms are the most important indicators of imminent risk; their disappearance may be associated with lower rates of transition to psychosis                                                                 |
| Stain et al 2016[141]       | Prevention psychosis     | The CHR-P criteria provide an important opportunity for early intervention in preventing or delaying the onset of psychosis and reducing the social and economic burden associated with long-term mental health problems                          |
| van der Gaag et al 2012[102; 142] | Prevention psychosis | Postponement or prevention of the transition to frank psychosis is the main goal: early detection is of little use without an effective intervention.                                                                                                               |
| Woods et al 2017[143; 144]  | Prevention psychosis     | To investigate the safety and efficacy of ziprasidone in delaying or preventing conversion to psychosis among individuals meeting CHR-P criteria                                                                                                                                  |
| Yung et al[145]; McGorry et al[146] 2011 | Prevention psychosis | Cognitive therapy and/or low-dose antipsychotic administered during the prodromal phase of schizophrenia may prevent or delay the onset of full-blown illness.                                                                                      |
| Trial title: ClinicalTrials.gov Identifier | Population (age); instrument | Trial arms | Allocation and masking | Duration | Sample size | Primary outcome |
|------------------------------------------|-----------------------------|------------|-----------------------|----------|-------------|----------------|
| The Role of Antidepressants or Antipsychotics in Preventing Psychosis: Fluoxetine vs Aripiprazole Comparative Trial (FACT). NCT02357849 | CHR-P (12-25); SIPS | 1. Fluoxetine; 2. aripiprazole | randomised (participant, care provider, investigator outcomes assessor) | 24 weeks | 48 | Time to either all-cause-discontinuation or need to add another psychotropic agent |
| Multimodal Prevention of Psychosis - Investigating Efficacy of N-Acetylcysteine and Psychotherapy in CHR-Patients (ESPRIT-B1). NCT03149107 | CHR-P (18-40); SIPS or SPI-A | 1. Integrated Preventive Psychological Intervention plus N-Acetylcysteine; 2. Psychological stress management and N-Acetylcysteine; 3. Integrated Preventive Psychological Intervention and placebo; 4. Psychological stress management and placebo | randomised (participant, investigator, outcomes assessor) | 6 months | 200 | Transition to psychosis |
| Randomized Controlled Trial of Aspirin vs Placebo in the Treatment of Patients With the Clinical Risk Syndrome for Psychosis. NCT02047539 | CHR-P (19-35); SIPS | 1. Aspirin (2-Acetoxybenzoic acid); 2. placebo | randomised (participant, care provider, investigator outcomes assessor) | 12 weeks? | 40 | Symptoms improvement |
| Placebo-controlled Trial in Subjects at Ultra-high Risk for Psychosis With Omega-3 Fatty Acids in Europe (PURPOSE). NCT02597439 | CHR-P (13-20); CAARMS | 1. Omega-3 fatty acids; 2. placebo | randomised (participant, care provider, investigator outcomes assessor) | 6 months | 220 | Transition to psychosis |
| Effects of Neurocognitive and Social Cognitive Remediation in Patients at Ultra-High Risk of Psychosis (FOCUS). NCT02098408 | CHR-P (18-40); CAARMS | 1. Standard treatment + cognitive remediation; 2. standard treatment | randomised (investigator, outcomes assessor) | 6 months | 126 | Cognitive functioning |
| Cognitive Behavioral Social Skills Training for Youth at Risk of Psychosis. NCT02234258 | CHR-P (14-30); SIPS | 1. Cognitive behavioral social skills; 2. psychoeducation | randomised (outcomes assessor) | 18 weeks | 225 | Social functioning |
| Decreasing Risk of Psychosis by Sulforaphane (DROPS Trial). NCT03932136 | CHR-P (15-45); SIPS | 1. Sulforaphane; 2. placebo | randomised (participant, care provider, investigator) | 52 weeks | 300 | Transition to psychosis |
| Study Title                                                                 | CHR-P (Age Range) | SIPS | Intervention                                                                 | Randomisation | Outcomes Assessor | Duration | Sample Size | Outcome Measures                  |
|----------------------------------------------------------------------------|-------------------|------|------------------------------------------------------------------------------|---------------|-------------------|----------|-------------|-----------------------------------|
| Targeted Cognitive Training in Clinical High Risk (CHR) for Psychosis. NCT02404194 | (15-30); SIPS     |      | 1. Targeted cognitive training; 2. computer game                              |               |                   | 10 weeks | 76          | Cognitive functioning             |
| Minocycline and/or Omega-3 Fatty Acids Added to Treatment as Usual for At Risk Mental States (NAYAB). NCT02569307 | (16-35); CAARMS   |      | 1. Minocycline; 2. omega-3 fatty acids; 3. treatment as usual                 |               |                   | 6 months | 320         | Transition to psychosis           |
| The Staged Treatment in Early Psychosis Study (STEP). NCT02751632           | (15-25); CAARMS   |      | staged treatment: 1. support and problem solving therapy; 2. cognitive behavioural case management; 3. cognitive behavioural case management plus fluoxetine; 4. cognitive behavioural case management plus placebo |               |                   | up to 12 months | 340        | Global functioning               |
| Exercise and Markers of Medial Temporal Health in Youth at Ultra High-risk for Psychosis. NCT02155699 | (16-24); SIPS     |      | 1. Exercise 1; 2. exercise 2; 3. waiting list                                |               |                   | 3 months | 45          | Brain volume                      |
| Glutamate Reducing Interventions in Schizophrenia. NCT03321617              | (18-30); NA       |      | 1. pomaglumetad methionil 40 mg, 2. pomaglumetad methionil 80 mg, 3. pomaglumetad methionil 120 mg, 4. pomaglumetad methionil 160 mg |               |                   | 2 weeks  | 50          | Cerebral blood volume             |
| Transcranial Direct Current Stimulation Coupled With Virtual Rehabilitation for Negative Symptoms in At-Risk Youth. NCT02951208 | (16-30); SIPS     |      | 1. Active Transcranial Direct Current Stimulation Coupled With Virtual Rehabilitation; 2. sham conditions |               |                   | 4 weeks  | 22          | Symptoms improvement              |
| Neurofeedback Processing Speed Training to Improve Social Functioning in Teenagers and Young Adults at Clinical High Risk for Psychosis. NCT03447548 | (12-25); SIPS     |      | 1. Neurofeedback processing speed training, 2. control                       |               |                   | NA       | 105         | Cognitive functioning             |
| Study Title                                                                 | DSM-5-APS          | Phase | Randomised | Duration | Participants | Primary Outcome |
|---------------------------------------------------------------------------|--------------------|-------|------------|----------|--------------|-----------------|
| A Phase II Randomised, Double-blind, Placebo-controlled Study to Evaluate the Efficacy, Safety, and Tolerability of Orally Administered BI 409306 During a 52-week Treatment Period as an Early Intervention in Patients With Attenuated Psychosis Syndrome. NCT03230097 | (16-30); DSM-5     | II     | participant, investigator | 52 weeks | 300            | Time to remission |
Figure 4. Example of sequential meta-analysis testing the preventive efficacy of CBT in CHR-P individuals. Plotted are the Randomised Controlled Trials of CBT vs Needs Based Intervention (NBI) in CHR-P patients that reported risk of psychosis onset at 6-month (part a) and 12 months (part b). The blue line represents the Z-value of each interim meta-analysis, the green line indicates the statistical significance threshold and the dotted red line the monitoring boundary. The red vertical line represents the a priori information size (APIS), i.e., the required sample size to detect a Relative Risk Reduction (RRR) = 0.5 with statistical power 0.9 on the risk of psychosis. Estimated using the package “metacumbounds[147]” and the previously published meta-analytical results[30].

a)
Figure 5. Clinical Stratification of Pretest Risk Enrichment in Individuals undergoing a Clinical High Risk assessment[113].
| Sampling | Recruitment (pretest) | Psychometric assessment (post-test) | Total sample size excluding attrition (c) |
|----------|----------------------|------------------------------------|--------------------------------------|
|          | Risk of psychosis at 3yrs | Risk of psychosis at 3yrs | Risks Ratio (Risk experimental treatment / Risk needs-based intervention) |
| Type of sample | CHR+ (a) | CHR- (b) | 0.5 | 0.6 | 0.7 |
| Pretest risk in people undergoing CHR assessment outside randomised clinical trials | 0.03(d)-0.49 | 0.051 | 0.003 | 2368 | 3942 | 7436 |
| General population | 0.004 | 0.007 | <0.001 | 17908 | 29844 | 56360 |
| Neurapro trial -control arm | 0.082 | 0.140 | 0.008 | 796 | 1324 | 2488 |
| Pretest risk stratification | | | | | | |
| low pretest risk | 0.014 | 0.026 | 0.001 | 4730 | 7880 | 14874 |
| moderately low pretest risk | 0.100 | 0.168 | 0.010 | 648 | 1076 | 2020 |
| moderately high pretest risk | 0.181 | 0.287 | 0.020 | 336 | 554 | 1036 |
| high pretest risk | 0.456 | 0.604 | 0.070 | 104 | 170 | 310 |

(a) LR+ = 1.82; (b) LR- = 0.09; (c) alpha=0.05; power 90%; 2-sided; allocation ratio = 1. Post-test probability=LR*pretest probability/[(1-pretest probability)+(pretest probability*LR)]. The sample sizes reported in the table indicate the individuals who should complete the trial (to estimate the baseline sample, attrition should be considered); (d) the average is 0.15 but because it depends on unstandardised idiosyncratic recruitment strategies, it is highly variable ranging from 3% to 49% depending on the study[5]. It tends to be on the lower side when the recruitment focuses on children and adolescent populations, intermediate when the recruitment focuses on primary care settings and higher when the recruitment focuses on secondary care. In this table to allow a conservative estimate of the sample size required we use only the lower bound of 0.03.
Currently, no reliable recommendation can be made regarding whether specific interventions (e.g. psychological interventions, medications, dietary interventions, needs-based interventions) are more effective compared to each other for the prevention of psychosis in CHR-P individuals.

Consequently, the safest approach is recommended, that is needs-based interventions and psychological interventions over antipsychotics, because the latter are not more efficacious than other options and have known side effects.

The selection of these two interventions should be based on factors such as the characteristics of each individual. These can include patients’ preferences (e.g., some patients may prefer psychological interventions over needs-based interventions), social circumstances (e.g. needs based interventions which include housing/vocational support may be suited for patients for whom these issues represent the presenting complaint), nature of symptoms (e.g. psychological interventions may be indicated for those presenting with cognitive biases in addition to attenuated psychotic symptoms), predicted risk (e.g. those presenting with brief and limited intermittent psychotic symptoms may need psychological treatments beyond needs-based interventions) or the local availability of each intervention.

Some suggestions can be made on the basis of differences that, even if non-statistically significant, had at least a moderate effect size (Odds Ratio > 2.5 for preventing the onset of psychosis, or Cohen’s d > 0.5 for reducing symptoms).

- The most efficacious intervention for preventing psychosis onset could be Integrated Psychological Interventions (IPI), and the second most efficacious could be Cognitive Behavioural Therapy (van der Gaag protocol) combined with needs-based interventions.

- Omega-3 and CBT could be more efficacious than IPI for improving attenuated positive psychotic symptoms.

- N-methyl-D-aspartate receptor modulators could be more efficacious than needs-based interventions alone, CBT and family therapy for improving attenuated negative symptoms.

- Importantly, none of these differences reached statistical significance.

Finally, it will be essential to consult the results of new and forthcoming studies as they emerge. In this regard, living meta-analyses could provide new evidence earlier than updating a conventional meta-analysis.
REFERENCES:

[1] P. Fusar-Poli, P.D. McGorry, and J.M. Kane, Improving outcomes of first-episode psychosis: an overview. World Psychiatry 16 (2017) 251-265.

[2] D. Oliver, J. Radua, A. Reichenberg, R. Uher, and P. Fusar-Poli, Psychosis Polyrisk Score (PPS) for the detection of individuals at risk and the prediction of their outcomes. Frontiers in Psychiatry (2019) in press.

[3] P. Fusar-Poli, S. Borgwardt, A. Bechdolf, J. Addington, A. Riecher-Rossler, F. Schultzze-Lutter, M. Keshavan, S. Wood, S. Ruhrmann, L.J. Seidman, L. Valsecchi, T. Cannon, E. Velthorst, L. De Haan, B. Comblatt, I. Bonoldi, M. Birchwood, T. McGlashan, W. Carpenter, P. McGorry, J. Klosterkotter, P. McGuire, and A. Yung, The psychosis high-risk state: a comprehensive state-of-the-art review. JAMA Psychiatry 70 (2013) 107-20.

[4] P. Fusar-Poli, M. Byrne, S. Badger, L.R. Valsecchi, and P.K. McGuire, Outreach and support in south London (OASIS), 2001-2011: ten years of early diagnosis and treatment for young individuals at high clinical risk for psychosis. Eur Psychiatry 28 (2013) 315-26.

[5] P. Fusar-Poli, M. Cappucciati, S. Borgwardt, S.W. Woods, J. Addington, B. Nelson, D.H. Nieman, D.R. Stahl, G. Rutigliano, A. Riecher-Rossler, A.E. Simon, M. Mizuno, T. Lee, J.S. Kwon, M.M.L. Lam, J. Perez, S. Keri, P. Amminger, S. Metzler, W. Kauhle, W. Roessler, J. Lee, J. Labad, T. Ziemans, S.K. An, C.-C. Liu, K.A. Woodberry, A. Brahm, C. Corcoran, P. McGorry, A.R. Yung, and P.K. McGuire, Heterogeneity of Psychosis Risk Within Individuals at Clinical High Risk A Meta-analytical Stratification. Jama Psychiatry 73 (2016) 113-120.

[6] J.R. Webb, J. Addington, D.O. Perkins, C.E. Bearden, K.S. Cadenhead, T.D. Cannon, B.A. Cornblatt, R.K. Heinsen, L.J. Seidman, S.I. Tarbox, M.T. Tsuang, E.F. Walker, T.H. McGlashan, and S.W. Woods, Specificity of Incident Diagnostic Outcomes in Patients at Clinical High Risk for Psychosis. Schizophr Bull 41 (2015) 1066-75.

[7] P. Fusar-Poli, Extending the Benefits of Indicated Prevention to Improve Outcomes of First-Episode Psychosis. JAMA Psychiatry 74 (2017) 667-668.
[16] I. Gartner, Gartner Hype Cycle, 2018.
[17] P. Fusar-Poli, The Hype Cycle of the Clinical High Risk State for Psychosis: the Need of a Refined Approach. Schizophr Bull (2018) EPub 6 January 2018.
[18] J.H. Chen, and S.M. Asch, Machine Learning and Prediction in Medicine - Beyond the Peak of Inflated Expectations. N Engl J Med 376 (2017) 2507-2509.
[19] P.D. McGorry, A.R. Yung, L.J. Phillips, H.P. Yuen, S. Fracey, E.M. Cosgrave, D. Germano, J. Bravin, T. McDonald, A. Blair, S. Adlard, and H. Jackson, Randomized Controlled Trial of Interventions Designed to Reduce the Risk of Progression to First-Episode Psychosis in a Clinical Sample With Subthreshold Symptoms. Archives of General Psychiatry 59 (2002) 921.
[20] T.H. McGlashan, R.B. Zipursky, D. Perkins, J. Addington, T. Miller, S.W. Woods, K.A. Hawkins, R.E. Hoffman, A. Preda, I. Epstein, D. Addington, S. Lindborg, Q. Trzaskoma, M. Tohen, and A. Breier, Randomized, double-blind trial of olanzapine versus placebo in patients prodromally symptomatic for psychosis. Am J Psychiatry 163 (2006) 790-799.
[21] A.P. Morrison, P. French, L. Walford, S.W. Lewis, A. Kilcommons, J. Green, S. Parker, and R.P. Bentall, Cognitive therapy for the prevention of psychosis in people at ultra-high risk: randomised controlled trial. Br J Psychiatry 185 (2004) 291-7.
[22] A. Bechdolf, M. Wagner, S. Ruhrmann, S. Harrigan, V. Puthoff, R. Pukrop, A. Brockhaus-Dumke, J. Berning, B. Janssen, P. Decker, R. Bottlender, K. Maurer, H.-J. Moller, W. Gaebel, H. Hafner, W. Maier, and J. Klosterkotter, Preventing progression to first-episode psychosis in early initial prodromal states. The British Journal of Psychiatry 200 (2012) 22-29.
[23] G.P. Amminger, M.R. Schäfer, K. Papageorgiou, M.C. Klier, M.S. Cotton, M.S. Harrigan, A. Mackinnon, P.D. McGorry, and E.G. Berger, Long-Chain omega-3 Fatty Acids for Indicated Prevention of Psychotic Disorders. Arch Gen Psychiatry 67 (2010) 146-154.
[24] NICE, Psychosis and schizophrenia in adults: prevention and management. National Institute for Clinical Excellence, https://www.nice.org.uk/Guidance/CG178, 2014.
[25] S.J. Schmidt, F. Schultze-Lutter, B.G. Schimmelmann, N.P. Maric, R.K.R. Salokangas, a. Riecher-Rössler, M. van der Gaag, a. Meneghelli, M. Nordentoft, M. Marshall, a. Morrison, a. Raballo, J. Klosterkötter, and S. Ruhrmann, EPA guidance on the early intervention in clinical high risk states of psychoses. European Psychiatry 30 (2015) 388-404.
[26] M.R. Stafford, H. Jackson, E. Mayo-Wilson, A.P. Morrison, and T. Kendall, Early interventions to prevent psychosis: systematic review and meta-analysis. BMJ 346 (2013) f185.
[27] A.P. Morrison, P. French, S.L. Stewart, M. Birchwood, D. Fowler, A.I. Gumley, P.B. Jones, R.P. Bentall, S.W. Lewis, G.K. Murray, P. Patterson, K. Brunet, J. Conroy, S. Parker, T. Reilly, R. Byrne, L.M. Davies, and G. Dunn, Early detection and intervention evaluation for people at risk of psychosis: multisite randomised controlled trial. BMJ 344 (2012) e2233.
[28] P.D. McGorry, B. Nelson, C. Markulev, H.P. Yuen, M.R. Schafer, N. Mossaheb, M. Schlegelhofer, S. Smesy, I.B. Hickie, G.E. Berger, E.Y. Shen, L. de Haan, D.H. Nieman, M. Nordentoft, A. Riecher-Rossler, S. Verma, A. Thompson, A.R. Yung, and G.P. Amminger, Effect of omega-3 Polyunsaturated Fatty Acids in Young People at Ultrahigh Risk for Psychotic Disorders: The NEURAPRO Randomized Clinical Trial. JAMA Psychiatry 74 (2017) 19-27.
[29] B. Nelson, G.P. Amminger, H.P. Yuen, C. Markulev, S. Lavoie, M.R. Schafer, J.A. Hartmann, N. Mossaheb, M. Schlegelhofer, S. Smesy, I.B. Hickie, G. Berger, E.Y.H. Chen, L. de Haan, D.H. Nieman, M. Nordentoft, A. Riecher-Rossler, S. Verma, A. Thompson, A.R. Yung, and P.D. McGorry, NEURAPRO: a multi-centre RCT of omega-3 polyunsaturated fatty acids versus placebo in young people at ultra-high risk of psychotic disorders-medium-term follow-up and clinical course. NPJ Schizophrenia 4 (2018) 11.
[30] C. Davies, A. Cipriani, J.P.A. Ioannidis, J. Radua, D. Stahl, U. Provenzani, P. McGuire, and P. Fusar-Poli, Lack of evidence to favor specific preventive interventions in psychosis: a network meta-analysis. World Psychiatry 17 (2018) 196-209.
[31] P. Fusar-Poli, Negative Psychosis Prevention Trials. JAMA Psychiatry (2017).
[32] B. Nelson, G.P. Amminger, and P.D. McGorry, Recent Meta-Analyses in the Clinical High Risk for Psychosis Population: Clinical Interpretation of Findings and Suggestions for Future Research. Front Psychiatry 9 (2018) 502.
[33] P. Fusar-Poli, and J. Radua, Ten simple rules for conducting Umbrella Reviews Evid Based Ment Health (2018) in press.
[34] R.D. Riley, P.C. Lambert, and G. Abo-Zaid, Meta-analysis of individual participant data: rationale, conduct, and reporting. BMJ 340 (2010) c221.
[35] B.J. Shea, C. Hamel, G.A. Wells, L.M. Bouter, E. Kristjansson, J. Grimshaw, D.A. Henry, and M. Boers, AMSTAR is a reliable and valid measurement tool to assess the methodological quality of systematic reviews. J Clin Epidemiol 62 (2009) 1013-20.
[36] A. Cipriani, X. Zhou, C. Del Giovane, S.E. Hetrick, B. Qin, C. Whittington, D. Coghill, Y. Zhang, P. Hazell, S. Leucht, P. Cuijpers, J. Pu, D. Cohen, A.V. Ravindran, Y. Liu, K.D. Michael, L. Yang, L. Liu, and P. Xie, Comparative efficacy and tolerability of antidepressants for major depressive disorder in children and adolescents: a network meta-analysis. The Lancet 388 (2016) 881-890.
[37] A.R. Jadad, D.J. Cook, A. Jones, T.P. Klassen, P. Tugwell, M. Moher, and D. Moher, Methodology and reports of systematic reviews and meta-analyses: a comparison of Cochrane reviews with articles published in paper-based journals. JAMA 280 (1998) 278-80.
[38] K.G. Shojania, M. Sampson, M.T. Ansari, J. Ji, S. Doucette, and D. Moher, How quickly do systematic reviews go out of date? A survival analysis. Annals of internal medicine 147 (2007) 224-33.
[39] A. Nikolakopoulou, D. Mavridis, T.A. Furukawa, A. Cipriani, A.C. Tricco, S.E. Straus, G.C.M. Siontis, M. Egger, and G. Salanti, Living network meta-analysis compared with pairwise meta-analysis in comparative effectiveness research: empirical study. BMJ 360 (2018) k585.
[40] P. Bragge, O. Clavisi, T. Turner, E. Tavender, A. Collie, and R.L. Gruen, The Global Evidence Mapping Initiative: scoping research in broad topic areas. BMC Med Res Methodol 11 (2011) 92.
[41] J.H. Elliott, T. Turner, O. Clavisi, J. Thomas, J.P. Higgins, C. Mavergames, and R.L. Gruen, Living systematic reviews: an emerging opportunity to narrow the evidence-practice gap. PLoS Med 11 (2014) e1001603.
[42] J.H. Elliott, A. Synnot, T. Turner, M. Simmonds, E.A. Akl, S. McDonald, G. Salanti, J. Meerpohl, H. MacLehose, J. Hilton, D. Tovey, I. Shemilt, J. Thomas, and N. Living Systematic Review, Living systematic review: 1. Introduction-the why, what, when, and how. J Clin Epidemiol 91 (2017) 23-30.
[43] K. Lan, and D. Demets, Discrete sequential boundaries for clinical trials. Biometrika 70 (1983) 659e63.
[44] J.M. Pogue, and S. Yusuf, Cumulating evidence from randomized trials: utilizing sequential monitoring boundaries for cumulative meta-analysis. Control Clin Trials 18 (1997) 580-93; discussion 661-6.
[45] J. Brok, K. Thorlund, J. Wetterslev, and C. Gluud, Apparently conclusive meta-analyses may be inconclusive--Trial sequential analysis adjustment of random error risk due to repetitive testing of accumulating data in apparently conclusive neonatal meta-analyses. Int J Epidemiol 38 (2009) 287-98.
[46] J. Wetterslev, K. Thorlund, J. Brok, and C. Gluud, Trial sequential analysis may establish when firm evidence is reached in cumulative meta-analysis. J Clin Epidemiol 61 (2008) 64-75.
[47] A. Nikolakopoulou, D. Mavridis, M. Egger, and G. Salanti, Continuously updated network meta-analysis and statistical monitoring for timely decision-making. Stat Methods Med Res 27 (2018) 1312-1330.
[48] J.A. Cook, J. Hislop, D.G. Altman, P. Fayers, A.H. Briggs, C.R. Ramsay, J.D. Norrie, I.M. Harvey, B. Buckley, D. Fergusson, I. Ford, L.D. Vale, and D. group, Specifying the target difference in the primary outcome for a randomised controlled trial: guidance for researchers. Trials 16 (2015) 12.
[49] M. Simmonds, G. Salanti, J. McKenzie, J. Elliott, and N. Living Systematic Review, Living systematic reviews: 3. Statistical methods for updating meta-analyses. J Clin Epidemiol 91 (2017) 38-46.

[50] J.P. Higgins, A. Whitehead, and M. Simmonds, Sequential methods for random-effects meta-analysis. Stat Med 30 (2011) 903-21.

[51] C. Michel, E. Toffel, S.J. Schmidt, S. Eliez, M. Armando, A. Solida-Tozzi, F. Schultz-Lutter, and M. Debbane, [Detection and early treatment of subjects at high risk of clinical psychosis: Definitions and recommendations]. Encephale 43 (2017) 292-297.

[52] M. van der Gaag, F. Smit, A. Bechdolf, P. French, D.H. Linszen, A.R. Yung, P. McGorry, and P. Cuijpers, Preventing a first episode of psychosis: meta-analysis of randomized controlled prevention trials of 12 month and longer-term follow-ups. Schizophr Res 149 (2013) 56-62.

[53] A. Preti, and M. Cella, Randomized-controlled trials in people at ultra high risk of psychosis: a review of treatment effectiveness. Schizophr Res 123 (2010) 30-6.

[54] C. Kelly, A.V. Hadjinicolaou, C. Holt, M. Agius, and R. Zaman, Meta-analysis of medical and non-medical treatments of the prodromal phase of psychotic illness in at-risk mental states. Psychiatria Danubina 22 (2010) 56-62.

[55] G. Deas, C. Kelly, A.V. Hadjinicolaou, and C. Holt, An update on: meta-analysis of medical and non-medical treatments of the prodromal phase of psychotic illness in at risk mental states. 28 (2016) 31-38.

[56] M. Marshall, and J. Rathbone, Early intervention for psychosis. in: M. Marshall, (Ed.), Cochrane Database of Systematic Reviews, John Wiley & Sons, Ltd, Chichester, 2011.

[57] P. Fusar-Poli, N. Werbeloff, G. Rutigliano, D. Oliver, C. Davies, D. Stahl, and P. McGuire, Transdiagnostic risk calculator for the automatic detection of individuals at risk and the prediction of psychosis: second replication in an independent national health service trust. Schizophr Bull (2018) EPub Jun 12 2018

[58] D.J. Devoe, M.S. Farris, P. Townes, and J. Addington, Attenuated psychotic symptom interventions in youth at risk of psychosis: A systematic review and meta-analysis. Early Interv Psychiatry (2018).

[59] M.R. Stafford, H. Jackson, E. Mayo-Wilson, A.P. Morrison, and T. Kendall, Errata: Early interventions to prevent psychosis: systematic review and meta-analysis. BMJ 346 (2013) f762-f762.

[60] J. Devoe, A. Peterson, and J. Addington, Erratum to: Devoe DJ, Peterson A, Addington J. Negative Symptom Interventions in Youth at Risk of Psychosis: A Systematic Review and Network Meta-Analys. Schizophr Bull 44 (2018) 463.

[61] J. Radua, V. Ramella-Cravaro, J.P.A. Ioannidis, A. Reichenberg, N. Phhiphophtatsanee, T. Amir, H. Yenn Thoo, D. Oliver, C. Davies, C. Morgan, P. McGuire, R.M. Murray, and P. Fusar-Poli, What causes psychosis? An umbrella review of risk and protective factors. World Psychiatry 17 (2018) 49-66.

[62] D.J. Devoe, A. Peterson, and J. Addington, Negative Symptom Interventions in Youth at Risk of Psychosis: A Systematic Review and Network Meta-analysis. Schizophr Bull 44 (2018) 807-823.

[63] C. Davies, J. Radua, A. Cipriani, D. Stahl, U. Provenzani, P. McGuire, and P. Fusar-Poli, Efficacy and Acceptability of Interventions for Attenuated Positive Psychotic Symptoms in Individuals at Clinical High Risk of Psychosis: A Network Meta-Analysis. Front Psychiatry 9 (2018) 187.

[64] P. Hutton, and P.J. Taylor, Cognitive behavioural therapy for psychosis prevention: a systematic review and meta-analysis. Psychol Med 44 (2014) 449-68.

[65] D.J. Devoe, M.S. Farris, P. Townes, and J. Addington, Interventions and social functioning in youth at risk of psychosis: A systematic review and meta-analysis. Early Interv Psychiatry (2018).

[66] J.T. Kantrowitz, S.W. Woods, E. Petkova, B. Cornblatt, C.M. Corcoran, H. Chen, G. Silipo, and D.C. Javitt, D-serine for the treatment of negative symptoms in individuals at clinical high risk of schizophrenia: A pilot, double-blind, placebo-controlled, randomised parallel group mechanistic proof-of-concept trial. The Lancet Psychiatry 2 (2015) 403-412.
D.J. Miklowitz, M.P. O'Brien, D.A. Schlosser, J. Addington, K.A. Candan, C. Marshall, I. Domingues, B.C. Walsh, J.L. Zinberg, S.D. De Silva, M. Friedman-Yakoobian, and T.D. Cannon, Family-Focused Treatment for Adolescents and Young Adults at High Risk for Psychosis: Results of a Randomized Trial. Journal of the American Academy of Child & Adolescent Psychiatry 53 (2014) 848-858.

S. Ruhrmann, A. Bechdolf, K.U. Kuhn, M. Wagner, F. Schulze-Lutter, B. Janssen, K. Maurer, H. Hafner, W. Gaebel, H.J. Moller, W. Maier, J. Klosterkotter, and L.s. group, Acute effects of treatment for prodromal symptoms for people putatively in a late initial prodromal state of psychosis. Br J Psychiatry Suppl 51 (2007) s88-95.

D. Piskulic, M. Barbato, L. Liu, and J. Addington, Pilot study of cognitive remediation therapy on cognition in young people at clinical high risk of psychosis. Psychiatry Res 225 (2015) 93-98.

R. Loewy, M. Fisher, D.A. Schlosser, B. Biagianti, B. Stuart, D.H. Mathalon, and S. Vinogradov, Intensive Auditory Cognitive Training Improves Verbal Memory in Adolescents and Young Adults at Clinical High Risk for Psychosis. Schizophr Bull 42 Suppl 1 (2016) S118-26.

J. Choi, C.M. Corcoran, J.M. Fiszdon, M. Stevens, D.C. Javitt, M. Deasy, L.C. Haber, M.J. Dewberry, and G.D. Pearlson, Pupilometer-based neurofeedback cognitive training to improve processing speed and social functioning in individuals at clinical high risk for psychosis. Psychiatr Rehabil J 40 (2017) 33-42.

M.F. Green, D.L. Penn, R. Bentall, W.T. Carpenter, W. Gaebel, R.C. Gur, A.M. Kring, S. Park, S.M. Silverstein, and R. Heinssen, Social cognition in schizophrenia: an NIMH workshop on definitions, assessment, and research opportunities. Schizophr Bull 34 (2008) 1211-20.

A.K. Fett, W. Viechtbauer, M.D. Dominguez, D.L. Penn, J. van Os, and L. Krabbendam, The relationship between neurocognition and social cognition with functional outcomes in schizophrenia: a meta-analysis. Neurosci Biobehav Rev 35 (2011) 573-88.

P. Fusar-Poli, G. Deste, R. Smieskova, S. Barlati, A.R. Yung, O. Howes, R.D. Stieglitz, A. Vita, P. McGuire, and S. Borgwardt, Cognitive functioning in prodromal psychosis: a meta-analysis. Arch Gen Psychiatry 69 (2012) 562-71.

R.J. van Donkersgoed, L. Wunderink, R. Nieboer, A. Aleman, and G.H. Pijnenborg, Social Cognition in Individuals at Ultra-High Risk for Psychosis: A Meta-Analysis. PLoS One 10 (2015) e0141075.

D. Feifel, and T. Reza, Oxytocin modulates psychotomimetic-induced deficits in sensorimotor gating. Psychopharmacology (Berl) 141 (1999) 93-8.

P.R. Lee, D.L. Brady, R.A. Shapiro, D.M. Dorsa, and J.I. Koenig, Social interaction deficits caused by chronic phencyclidine administration are reversed by oxytocin. Neuropsychopharmacology 30 (2005) 1883-94.

M. Kosfeld, M. Heinrichs, P.J. Zak, U. Fischbacher, and E. Fehr, Oxytocin increases trust in humans. Nature 435 (2005) 673-6.

M. Eckstein, B. Becker, D. Scheele, C. Scholz, K. Preckel, T.E. Schlaepfer, V. Grinevich, K.M. Kendrick, W. Maier, and R. Hurlemann, Oxytocin facilitates the extinction of conditioned fear in humans. Biol Psychiatry 78 (2015) 194-202.

C.F. Zink, and A. Meyer-Lindenberg, Human neuroimaging of oxytocin and vasopressin in social cognition. Horm Behav 61 (2012) 400-9.

C.A. Pedersen, C.M. Gibson, S.W. Rau, K. Salimi, K.L. Smedley, R.L. Casey, J. Leserman, L.F. Jaruskog, and D.L. Penn, Intranasal oxytocin reduces psychotic symptoms and improves Theory of Mind and social perception in schizophrenia. Schizophr Res 132 (2011) 50-3.

S.G. Hofmann, A. Fang, and D.N. Brager, Removal notice to Effect of Intranasal Oxytocin Administration on Psychiatric Symptoms: A Meta-Analysis of Placebo-Controlled Studies Psychiatr Res. 228 (2015) 708-714. Psychiatry Res 263 (2018) 299.

D.R. Williams, and P.C. Burkner, Effects of intranasal oxytocin on symptoms of schizophrenia: A multivariate Bayesian meta-analysis. Psychoneuroendocrinology 75 (2017) 141-151.
[84] R.A. Bethlehem, J. van Honk, B. Auyeung, and S. Baron-Cohen, Oxytocin, brain physiology, and functional connectivity: a review of intranasal oxytocin fMRI studies. Psychoneuroendocrinology 38 (2013) 962-74.

[85] S.A. Grace, S.L. Rossell, M. Heinrichs, C. Kordsachia, and I. Labuschagne, Oxytocin and brain activity in humans: A systematic review and coordinate-based meta-analysis of functional MRI studies. Psychoneuroendocrinology 96 (2018) 6-24.

[86] Y. Paloyelis, O.M. Doyle, F.O. Zelaya, S. Maltezos, S.C. Williams, A. Fotopoulou, and M.A. Howard, A Spatiotemporal Profile of In Vivo Cerebral Blood Flow Changes Following Intranasal Oxytocin in Humans. Biol Psychiatry 79 (2016) 693-705.

[87] C. Davies, Y. Paloyelis, G. Rutigliano, M. Cappucciati, A. De Micheli, V. Ramella-Cravaro, U. Provenzani, M. Antoniades, G. Modinos, D. Oliver, D. Stahl, S. Murgia, F. Zelaya, P. Allen, S. Shergill, P. Morrison, S. Williams, D. Taylor, P. McGuire, and P. Fusar-Poli, Oxytocin modulates hippocampal perfusion in people at clinical high risk for psychosis. Neuropsychopharmacology (2019).

[88] P. McGuire, P. Robson, W.J. Cubala, D. Vasile, P.D. Morrison, R. Barron, A. Taylor, and S. Wright, Cannabidiol (CBD) as an Adjunctive Therapy in Schizophrenia: A Multicenter Randomized Controlled Trial. Am J Psychiatry 175 (2018) 225-231.

[89] F.M. Leweke, D. Piomelli, F. Pahlisch, D. Muhl, C.W. Gerth, C. Hoyer, J. Klosterkotter, M. Hellmich, and D. Koethe, Cannabidiol enhances anandamide signaling and alleviates psychotic symptoms of schizophrenia. Transl Psychiatry 2 (2012) e94.

[90] S. Bhattacharyya, R. Wilson, E. Appiah-Kusi, A. O'Neill, M. Brammer, J. Perez, R. Murray, P. Allen, M.G. Bossong, and P. McGuire, Effect of Cannabidiol on Medial Temporal, Midbrain, and Strial Dysfunction in People at Clinical High Risk of Psychosis: A Randomized Clinical Trial. JAMA Psychiatry 75 (2018) 1107-1117.

[91] P. Fusar-Poli, G. Rutigliano, D. Stahl, C. Davies, I. Bonoldi, T. Reilly, and P. McGuire, Development and validation of a clinically based risk calculator for the transdiagnostic prediction of psychosis. Jama Psychiatry 74 (2017) 493-500.

[92] P.D. McGorry, J.A. Hartmann, R. Spooner, and B. Nelson, Beyond the "at risk mental state" concept: transitioning to transdiagnostic psychiatry. World Psychiatry 17 (2018) 133-142.

[93] P. Fusar-Poli, D. Oliver, G. Spada, R. Patel, R. Stewart, R. Dobson, and P. McGuire, Real-world Implementation of a Transdiagnostic Risk Calculator for the Automatic Detection of Individuals at Risk of Psychosis in Clinical Routine: Study Protocol. Frontiers Psychiatry (2019).

[94] M. McDonald, E. Christoforidou, N. Van Rijlsbergen, R. Gajwani, J. Gross, A.I. Gumley, S.M. Lawrie, M. Schwannauer, F. Schultz-Lutter, and P.J. Uhlhaas, Using Online Screening in the General Population to Detect Participants at Clinical High-Risk for Psychosis. Schizophr Bull (2018).

[95] M.J. Millan, A. Andrieux, G. Bartzokis, K. Cadenhead, P. Dazzan, P. Fusar-Poli, J. Gallinat, J. Giedd, D.R. Grayson, M. Heinrichs, R. Kahn, M.O. Krebs, M. Leboyer, D. Lewis, O. Marin, P. Marin, A. Meyer-Lindenberg, P. McGorry, P. McGuire, M.J. Owen, P. Patterson, A. Sawa, M. Spedding, P. Uhlhaas, F. Vaccarino, C. Wahlestedt, and D. Weinberger, Altering the course of schizophrenia: progress and perspectives. Nat Rev Drug Discov 5 (2016) 485-51.

[96] C. Davies, A. Cipriani, J. Radua, U. Provenzani, P. McGuire, and P. Fusar-Poli, Preventative treatments for psychosis: a network meta-analysis protocol. PROSPERO International prospective register of systematic reviews (2017).

[97] J.A. Hartmann, P.D. McGorry, S.J. Schmidt, G.P. Amminger, H.P. Yuen, C. Markulev, G.E. Berger, E.Y.H. Chen, L. de Haan, I.B. Hickie, S. Lavoie, M.J. McHugh, N. Mossaheb, D.H. Nieman, M. Nordentoft, A. Riecher-Rössler, M.R. Schäfer, M. Schlögelhofer, S. Smesny, A. Thompson, S.K. Verma, A.R. Yung, and B. Nelson, Opening the Black Box of Cognitive-Behavioural Case Management in Clients with Ultra-High Risk for Psychosis. Psychotherapy and Psychosomatics 3052 (2017) 292-299.

[98] A. Cipriani, J.P.T. Higgins, J.R. Geddes, and G. Salanti, Conceptual and technical challenges in network meta-analysis. Annals of internal medicine 159 (2013) 130-137.
[99] K. Thorlund, and E.J. Mills, Sample size and power considerations in network meta-analysis. Systematic Reviews 1 (2012).

[100] S. Kanters, N. Ford, E. Druyts, K. Thorlund, E.J. Mills, and N. Bansback, Use of network meta-analysis in clinical guidelines. Bulletin of the World Health Organization 94 (2016) 782-784.

[101] S. Leucht, A. Chaimani, A.S. Cipriani, J.M. Davis, T.A. Furukawa, and G. Salanti, Network meta-analyses should be the highest level of evidence in treatment guidelines. European Archives of Psychiatry and Clinical Neuroscience 266 (2016) 477-480.

[102] M. van der Gaag, D.H. Nieman, J. Rietdijk, S. Dragt, H.K. Ising, R.M.C. Klaassen, M. Koeter, P. Cuijpers, L. Wunderink, and D.H. Linszen, Cognitive Behavioral Therapy for Subjects at Ultrahigh Risk for Developing Psychosis: A Randomized Controlled Clinical Trial. Schizophrenia Bulletin 38 (2012) 1180-1188.

[103] P. Fusar-Poli, and J. Van Os, Lost in transition: setting the psychosis threshold in prodromal research. Acta Psychiatrica Scandinavica 127 (2013) 248-252.

[104] P. Fusar-Poli, M. Cappucciati, A. De Micheli, G. Rutigliano, I. Bonoldi, S. Tognin, V. Ramella-Cravaro, A. Castagnini, and P. McGuire, Diagnostic and Prognostic Significance of Brief Limited Intermittent Psychotic Symptoms (BLIPS) in Individuals at Ultra High Risk. Schizophr Bull 43 (2017) 47-56.

[105] P. Fusar-Poli, M. Cappucciati, I. Bonoldi, L.M. Hui, G. Rutigliano, D.R. Stahl, S. Borgwardt, P. Politi, A.L. Mishara, S.M. Lawrie, W.T. Carpenter, Jr., and P.K. McGuire, Prognosis of Brief Psychotic Episodes: A Meta-analysis. JAMA Psychiatry 73 (2016) 211-20.

[106] E.P. Moll van Charante, E. Richard, L.S. Eurelings, J.W. van Dalen, S.A. Ligthart, E.F. van Bussel, M.P. Hoevenaar-Blom, M. Vermeulen, and W.A. van Gool, Effectiveness of a 6-year multidomain vascular care intervention to prevent dementia (preDIVA): a cluster-randomised controlled trial. Lancet 388 (2016) 797-805.

[107] H. Frankish, and R. Horton, Prevention and management of dementia: a priority for public health. Lancet 390 (2017) 2614-2615.

[108] D. Oliver, C. Davies, G. Crossland, S. Lim, G. Gifford, P. McGuire, and P. Fusar-Poli, Can We Reduce the Duration of Untreated Psychosis? A Systematic Review and Meta-Analysis of Controlled Interventional Studies. Schizophr Bull (2018).

[109] D.G. Altman, and J.M. Bland, Absence of evidence is not evidence of absence. BMJ 311 (1995) 485.

[110] J. Cuzick, Preventive therapy for cancer. Lancet Oncol 18 (2017) e472-e482.

[111] P. Fusar-Poli, and F. Schultzze-Lutter, Predicting the onset of psychosis in patients at clinical high risk: practical guide to probabilistic prognostic reasoning. Evid Based Ment Health 19 (2016) 10-5.

[112] P. Fusar-Poli, Why ultra high risk criteria for psychosis prediction do not work well outside clinical samples and what to do about it. World Psychiatry 16 (2017) 212-213.

[113] P. Fusar-Poli, G. Rutigliano, D. Stahl, A. Schmidt, V. Ramella-Cravaro, H. Shetty, and P. McGuire, Deconstructing pretest risk enrichment to optimize prediction of psychosis in individuals at clinical high risk. Jama Psychiatry 73 (2016) 1260-1267.

[114] P. Fusar-Poli, F. Schultze-Lutter, and J. Addington, Intensive community outreach for those at ultra high risk of psychosis: dilution, not solution. Lancet Psychiatry 3 (2016) 18.

[115] P. Fusar-Poli, I. Bonoldi, A.R. Yung, S. Borgwardt, M.J. Kempton, L. Valmaggia, F. Barale, E. Caverzasi, and P. McGuire, Predicting psychosis: meta-analysis of transition outcomes in individuals at high clinical risk. Arch Gen Psychiatry 69 (2012) 220-9.

[116] P. Fusar-Poli, E. Palombini, C. Davies, D. Oliver, I. Bonoldi, V. Ramella-Cravaro, and P. McGuire, Why transition risk to psychosis is not declining at the OASIS ultra high risk service: The hidden role of stable pretest risk enrichment. Schizophr Res (2017).

[117] G. Gifford, N. Crossley, P. Fusar-Poli, H.G. Schnack, R.S. Kahn, N. Koutsouleris, T.D. Cannon, and P. McGuire, Using neuroimaging to help predict the onset of psychosis. Neuroimage 145 (2017) 209-217.

[118] P. Fusar-Poli, D. Stringer, A. Durieux, G. Rutigliano, I. Bonoldi, A. De Micheli, and D. Stahl, Clinical-learning versus machine learning for transdiagnostic prediction of psychosis onset in individuals at-risk Transl Psychiatry (2019) in press.
[119] A. Schmidt, M. Cappucciati, J. Radua, G. Rutigliano, M. Rocchetti, L. Dell’Osso, P. Politi, S. Borgwardt, T. Reilly, L. Valmaggia, P. McGuire, and P. Fusar-Poli, Improving Prognostic Accuracy in Subjects at Clinical High Risk for Psychosis: Systematic Review of Predictive Models and Meta-analytical Sequential Testing Simulation. Schizophr Bull 43 (2017) 375-388.

[120] J.M. Kane, and C.U. Correll, omega-3 Polyunsaturated Fatty Acids to Prevent Psychosis: The Importance of Replication Studies. JAMA Psychiatry 74 (2017) 11-12.

[121] G. Rutigliano, S. Merlino, A. Minichino, R. Patel, C. Davies, D. Oliver, A. De Micheli, P. McGuire, and P. Fusar-Poli, Long term outcomes of acute and transient psychotic disorders: The missed opportunity of preventive interventions. Eur Psychiatry 52 (2018) 126-133.

[122] A. Minichino, G. Rutigliano, S. Merlino, C. Davies, D. Oliver, A. De Micheli, R. Patel, P. McGuire, and P. Fusar-Poli, Unmet needs in patients with brief psychotic disorders: too ill for clinical high risk services and not enough ill for first episode services. European Psychiatry (2018).

[123] T.D. Cannon, C. Yu, J. Addington, C.E. Bearden, K.S. Cadenhead, B.A. Comblatt, R. Heinssen, C.D. Jeffries, D.H. Mathalon, T.H. McGlashan, D.O. Perkins, L.J. Seidman, M.T. Tsuang, E.F. Walker, S.W. Woods, and M.W. Kattan, An Individualized Risk Calculator for Research in Prodromal Psychosis. Am J Psychiatry (2016) appiajp201615070890.

[124] N. Koutsouleris, L. Kambeitz-Illankovic, S. Ruhrmann, M. Rosen, A. Ruef, D.B. Dwyer, M. Paolini, K. Chisholm, J. Kambeitz, T. Haidl, A. Schmidt, J. Gillam, F. Schultz-Lutter, P. Falkai, M. Reiser, A. Riecher-Rossler, R. Upthegrove, J. Hietala, R.K.R. Salokangas, C. Panetis, E. Meisenzahl, S.J. Wood, D. Beque, P. Brambilla, S. Borgwardt, and P. Consortium, Prediction Models of Functional Outcomes for Individuals in the Clinical High-Risk State for Psychosis or With Recent-Onset Depression: A Multimodal, Multisite Machine Learning Analysis. JAMA Psychiatry 75 (2018) 1156-1172.

[125] R. Uher, and A. Zwicker, Etiology in psychiatry: embracing the reality of poly-gene-environmental causation of mental illness. World Psychiatry 16 (2017) 121-129.

[126] J. Thomas, A. Noel-Storr, I. Marshall, B. Wallace, S. McDonald, C. Mavergames, P. Glasziou, I. Shemilt, A. Synnot, T. Turner, J. Elliott, and N. Living Systematic Review, Living systematic reviews: 2. Combining human and machine effort. J Clin Epidemiol 91 (2017) 31-37.

[127] D. O'Brien, K. Harvey, J. Howse, T. Reardon, and C. Creswell, Barriers to managing child and adolescent mental health problems: a systematic review of primary care practitioners' perceptions. Br J Gen Pract 66 (2016) e693-707.

[128] HM Government, No health without mental health: implementation framework. in: https://www.gov.uk/government/uploads/system/uploads/attachment_data/file/215811/, and d.p.a.S. 2016). (Eds.), 2011.

[129] NHS England, Future in mind: promoting, protecting and improving our children and young people’s mental health and wellbeing. in: Department of Health, (Ed.), 2015.

[130] C.G. Davey, and P.D. McGorry, Early intervention for depression in young people: a blind spot in mental health care. Lancet Psychiatry (2018).

[131] E. Vieta, E. Salagre, I. Grande, A.F. Carvalho, B.S. Fernandes, M. Berk, B. Birmaher, M. Tohen, and T. Suppes, Early Intervention in Bipolar Disorder. Am J Psychiatry 175 (2018) 411-426.

[132] P. Fusar-Poli, A. De Micheli, M. Rocchetti, M. Cappucciati, V. Ramella-Cravaro, G. Rutigliano, I. Bonoldi, P. McGuire, and I. Falkenberg, Semistructured Interview for Bipolar At Risk States (SIBARS). Psychiatry Res 264 (2018) 302-309.

[133] P.D. McGorry, and C. Mei, Early intervention in youth mental health: progress and future directions. Evid Based Ment Health 21 (2018) 182-184.

[134] P.D. McGorry, S.D. Goldstone, A.G. Parker, D.J. Rickwood, and I.B. Hickie, Cultures for mental health care of young people: an Australian blueprint for reform. Lancet Psychiatry 1 (2014) 559-68.

[135] S.E. Hetrick, A.P. Bailey, K.E. Smith, A. Malla, S. Mathias, S.P. Singh, A. O'Reilly, S.K. Verma, L. Benoit, T.M. Fleming, M.R. Moro, D.J. Rickwood, J. Duffy, T. Eriksen, R. Illback,
C.A. Fisher, and P.D. McGorry, Integrated (one-stop shop) youth health care: best available evidence and future directions. Med J Aust 207 (2017) S5-S18.

[136] C.C. Hagan, J.M. Graham, P.O. Wilkinson, N. Midgley, J. Suckling, B.J. Sahakian, and I.M. Goodyer, Neurodevelopment and ages of onset in depressive disorders. Lancet Psychiatry 2 (2015) 1112-6.

[137] ClinicalTrials.gov, The Staged Treatment in Early Psychosis Study (STEP), U.S. National Library of Medicine, 2019.

[138] J. Addington, I. Epstein, L. Liu, P. French, K.M. Boydell, and R.B. Zipursky, A randomized controlled trial of cognitive behavioral therapy for individuals at clinical high risk of psychosis. Schizophrenia Research 125 (2011) 54-61.

[139] A. Bechdolf, H. Müller, H. Stützer, M. Lambert, A. Karow, M. Zink, M. Lautenschlager, A. Heinz, W. de Millas, B. Janssen, W. Gaebel, F. Schneider, G. Juckel, S. Krüger-Özgürdal, T. Wobrock, M. Wagner, W. Maier, and J. Klosterkötter, 108. PREVENT: A Randomized Controlled Trial for the Prevention of First-Episode Psychosis Comparing Cognitive–Behavior Therapy (CBT), Clinical Management, and Aripiprazole Combined and Clinical Management and Placebo Combined. Schizophrenia Bulletin 43 (2016) S56.

[140] K. Cadenhead, J. Addington, T. Cannon, B. Cornblatt, D. Mathalon, T. McGlashan, D. Perkins, L.J. Seidman, M. Tsuang, E. Walker, and S. Woods, 23. Omega-3 Fatty Acid Versus Placebo in a Clinical High-Risk Sample From the North American Prodrome Longitudinal Studies (NAPLS) Consortium. Schizophrenia Bulletin 43 (2017) S16-S16.

[141] H.J. Stain, S. Bucci, A.L. Baker, V. Carr, R. Emsley, S. Halpin, T. Lewin, U. Schall, V. Clarke, K. Crittenden, and M. Startup, A randomised controlled trial of cognitive behaviour therapy versus non-directive reflective listening for young people at ultra high risk of developing psychosis: The detection and evaluation of psychological therapy (DEPTh) trial. Schizophrenia Research 176 (2016) 212-219.

[142] H.K. Ising, T.C. Kraan, J. Riedtijk, S. Dragt, R.M.C. Klaassen, N. Boonstra, D.H. Nieman, M. Willebrands-Mendrik, D.P.G. van den Berg, D.H. Linszen, L. Wunderink, W. Veling, F. Smit, and M. van der Gaag, Four-Year Follow-up of Cognitive Behavioral Therapy in Persons at Ultra-High Risk for Developing Psychosis: The Dutch Early Detection Intervention Evaluation (EDIE-NL) Trial. Schizophrenia Bulletin 42 (2016) 1243-52.

[143] S. Woods, J. Saksa, M. Compton, M. Daley, R. Rajarethinam, K. Graham, N. Breitborde, J. Cahill, V. Srihari, D. Perkins, C. Bearden, T. Cannon, E. Walker, and T. McGlashan, 112. Effects of Ziprasidone Versus Placebo in Patients at Clinical High Risk for Psychosis. Schizophrenia Bulletin 43 (2017) S58-S58.

[144] S.W. Woods, Ziprasidone in the Psychosis Prodrome (ZIP) ClinicalTrials.gov Identifier: NCT00635700 [Last updated: September 6, 2016]. ClinicalTrials.gov (2016).

[145] A.R. Yung, L.J. Phillips, B. Nelson, S.M. Francye, H. Pan/Yuen, M.B. Simmons, M.L. Ross, D. Kelly, K. Baker, G.P. Amminger, G. Berger, A.D. Thompson, A. Thampi, and P.D. McGorry, Randomized Controlled Trial of Interventions for Young People at Ultra High Risk for Psychosis. The Journal of Clinical Psychiatry 72 (2011) 430-440.

[146] P.D. McGorry, B. Nelson, L.J. Phillips, H.P. Yuen, S.M. Francye, A. Thampi, G.E. Berger, P. Amminger, M.B. Simmons, D. Kelly, A.D. Thompson, and A.R. Yung, Randomized controlled trial of interventions for young people at ultra-high risk of psychosis: Twelve-month outcome. Journal of Clinical Psychiatry 74 (2013) 349-356.

[147] B. Miladinovic, I. Hozo, and B. Djulbegovic, Trial sequential boundaries for cumulative meta-analyses. The Stata Journal 13 (2013) 77-91.