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Sleep abnormalities in different clinical stages of psychosis: A systematic review and meta-analysis

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33 Key points

• **Question**: Do sleep abnormalities differ in occurrence and severity in clinical high-risk (CHR-P), early psychosis (EP), and chronic psychosis (CP)? Findings: Sleep disturbance prevalence across 5135 cases was 50% and was comparable • across psychosis stages. Comparing 1575 cases and 977 controls revealed poor self-reported sleep quality throughout stages. CP had more arousal vs. CHR-P and reduced spindle duration vs. EP. Meaning: These findings indicate that a) sleep disturbances are highly prevalent • throughout psychosis stages; b) CHR-P, EP, and CP show common and distinct self-reported and objective sleep alterations, thus representing clinical targets and research domains for psychosis.

56	Abstract

57 Importance: Abnormal sleep is frequent in psychosis; however, sleep abnormalities in different
58 stages (i.e., Clinical-High-risk for Psychosis (CHR-P), Early Psychosis (EP), and Chronic
59 Psychosis (CP)) have not been characterized.

60 **Objective**: Identify sleep abnormalities across psychosis stages.

Data sources: Web of Science and PubMed were searched between inception and June 15th,
2022.

63 Study selection: Sleep disturbance prevalence studies and case-control studies reporting sleep
64 quality, sleep architecture, or sleep EEG oscillations in CHR-P, EP, or CP.

Data Extraction and Synthesis: This meta-analysis (PROSPERO; <u>CRD42021240503</u>) followed PRISMA 2020 guidelines. Stage-specific and pooled random-effects meta-analyses were conducted, along with the assessment of heterogeneity, study quality, and meta-regressions (clinical stage, sex, age, medication status, psychotic symptoms).

Main Outcomes and Measures: Sleep disturbance prevalence, self-reported sleep quality, sleep architecture (total sleep time, sleep latency, sleep efficiency, NREM and REM stages, number of arousals), and sleep EEG oscillations (spindle density, amplitude, and duration, and slow wave density).

Results: Fifty-nine studies with up to 6710 cases (N= 5135 for prevalence) and 977 controls were included. Sleep disturbance prevalence in pooled cases was 50% (95%CI=40-61%) and it was similar in each psychosis stage. Sleep quality was worse in pooled cases vs. controls (standardized mean difference, SMD=1.00, 95%CI=[0.70-1.30]). Sleep architecture alterations included: higher sleep onset latency (pooled cases SMD=0.96[0.62-1.30], EP SMD=0.72[0.52-

78 0.92], CP SMD=1.36[0.66-2.05]), higher wake after sleep onset (pooled cases SMD=0.5[0.29-79 0.71], EP SMD=0.62[0.34-0.89], CP SMD=0.51[0.09-0.93]), higher number of arousals (pooled 80 cases SMD=0.45[0.07-0.83], CP SMD=0.81[0.30-1.32]), higher stage 1 sleep (pooled cases 81 SMD=0.23[0.06-0.40], EP SMD=0.34[0.15-0.53]), lower sleep efficiency (pooled cases SMD=-82 0.75[-0.98 to -0.52], EP SMD=-0.90[-1.20 to -0.60], CP SMD=-0.73[-1.14 to -0.33]), and lower 83 rapid eye movement density (pooled cases SMD=0.37[0.14-0.60], CP SMD=0.48[0.19-0.77]). 84 Spindle parameter deficits included density: pooled cases SMD=-1.06[-1.50 to -0.63], EP 85 SMD=-0.80[-1.22 to -0.39], CP SMD=-1.39[-2.05 to -0.74]; amplitude: pooled cases SMD=-86 1.08[-1.33 to -0.82], EP SMD=-0.86[-1.24 to -0.47], CP SMD=-1.25[-1.58 to -0.91]; and 87 duration: pooled cases SMD=-1.21[-1.69 to -0.73], EP SMD=-0.71[-1.08 to -0.34], CP SMD=-88 1.74[-2.10 to -1.38]. Furthermore, CP had more frequent arousals vs. CHR-P (z=2.24, p=0.02), 89 and reduced spindle duration vs EP (z=-3.91, p<0.001). 90 Conclusion: Sleep disturbances are highly prevalent throughout the course of psychosis, and

different psychosis stages show both shared and distinct abnormalities in sleep quality,
architecture, and spindles. Thus, sleep should become a core clinical target and research domain
from at-risk to early and chronic stages of psychosis.

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102 Introduction

Sleep abnormalities have been observed in psychotic disorders since the dawn of psychiatric literature¹. Sleep disturbances, such as insomnia, are commonly reported by individuals with chronic psychosis $(CP)^2$ and are associated with subsequent relapse³. Altered sleep often precedes a psychotic episode in early psychosis $(EP)^4$, and disrupted sleep contributes to predicting transition to psychosis in youth at clinical high risk $(CHR-P)^5$. Thus, sleep abnormalities not only co-occur with psychotic symptoms but are also implicated in the development, manifestation, and recurrence of psychosis⁶.

Sleep disturbance prevalence, which is usually assessed with self-reported questionnaires (e.g., the Pittsburgh Sleep Quality Index, PSQI)⁷ is ~25% in the general population^{7,8}. Several studies have reported higher sleep disturbance prevalence in psychosis, although rates vary substantially (21-100%)⁹⁻¹¹ and have thus far never been meta-analyzed in different psychosis stages.

114 Altered sleep patterns across psychosis stages can also be examined in case control comparisons. 115 Several case control studies have used subjective sleep assessments (e.g., PSQI), which are 116 inexpensive and easy to implement in large clinical cohorts, and have reported worse sleep quality in CHR-P¹², EP¹³, and CP¹⁴ vs. healthy comparison groups. Other sleep studies have 117 118 utilized actigraphy, electroencephalography (EEG), and polysomnography (PSG) to objectively 119 quantify altered sleep characteristics in psychosis. Traditionally, these studies have focused on sleep architecture. Meta-analyses of sleep architecture findings from actigraphy¹⁵ and PSG/EEG 120 studies^{16–18} revealed shorter total sleep time and longer sleep onset latency and wake after sleep 121

onset in CP. PSG/EEG studies also showed decreased deep NREM sleep and reduced latency and duration of REM sleep in these patients¹⁸. Furthermore, shorter total sleep time and larger wake after sleep onset were reported in EP and CHR-P¹⁹, suggesting that altered sleep architecture is an early feature of psychosis.

More recently, several studies investigated sleep-specific EEG oscillations, including spindles, in psychosis. Deficits in spindle parameters (i.e., density, amplitude, and duration) were established in CP^{20} and $EP^{21,22}$. Furthermore, a recent meta-analysis reported reduced spindle density in psychotic disorders that yielded large effect sizes and was associated with disease progression²³.

Systematically investigating the occurrence and severity of sleep abnormalities in CHR-P, EP, and CP may therefore help differentiate sleep dysfunctions associated with chronicity and longterm medication exposure (i.e., observed only/primarily in CP) from those implicated in the manifestation of full-blown psychosis (i.e., occurring first in EP) and from sleep alterations related to vulnerability to psychosis (i.e., present since CHR-P)²⁴. This meta-analysis assessed, for the first time, the prevalence of sleep disturbances, along with subjective and objective sleep alterations throughout the course of psychosis, including CHR-P, EP, and CP.

137 Methods

This meta-analysis followed the Preferred Reporting Items for Systematic Reviews and Meta analyses (PRISMA) 2020²⁵ guidelines. The protocol was registered in PROSPERO
 (CRD42021240503).

141 Inclusion and exclusion criteria

For inclusion, studies had to be published between inception and June 15th, 2022, and written in English. Diagnosis of stages of psychosis was established using a recognized clinical assessment tool (see eMethods for *Clinical Stages* definition). Studies needed to provide measures of the prevalence of sleep disturbances in individuals at different psychosis stages and/or quantification of sleep characteristics in these individuals, assessed with PSG, EEG, actigraphy, or self-reports. Inclusion and exclusion criteria are explained in greater detail in the eMethods.

148 *Search strategy*

One author (JB) performed the literature search on the Web of Science and PubMed from inception until June 15th, 2022. The description of study search terms is provided in the eMethods. A manual search of the references of included articles and of relevant prior reviews/meta-analyses were also performed.

153 The eMethods contain details on study selection and data extraction.

154 Methodological quality appraisal

Quality appraisal was assessed using the Agency for Healthcare Research and Quality (AHRQ)²⁶
methodology checklist for cross-sectional/prevalence studies. For additional details, see
eMethods.

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160 Statistical analysis

Sleep disturbance prevalence was evaluated in three distinct analyses: 1) a pooled cases analysis of sleep disturbance prevalence aggregating all psychosis stages; 2) a stage-specific cases analysis of sleep disturbance; and 3) a moderator analysis comparing clinical stages with one another. We also performed a secondary analysis in a subgroup of studies assessing insomnia. Sleep disturbance prevalence effect sizes were analyzed as logit transformed values, quantifying the log odds of sleep disturbance.

167 Sleep architecture and oscillatory alterations were evaluated in three different analyses: 1) a 168 pooled case-control comparison of each sleep variable aggregating all psychosis stages; 2) a 169 stage-specific case-control analysis of sleep abnormalities; and 3) a moderator analysis 170 comparing clinical stages with one another. Sleep architecture and sleep oscillatory parameters were analyzed as standardized mean differences across groups using the Hedges' g statistic²⁷. 171 172 For all hypothesis testing, we used two-sided tests with statistical significance at the P<.05 level. 173 A random effects linear regression model was fitted for each sleep parameter, and calculated 174 effect sizes were weighted according to inverse variance to account for the variability of each study²⁸. Meta-analysis models were estimated using restricted maximum likelihood estimation 175 176 using the *rma* function in the R *metafor* package in R software v. 4.1.0 (method = "PLO" for prevalence analyses; method = "SMD" for case-control comparisons). 177

The recovery of missing or partial data from studies and the assessment of study heterogeneity
using funnel plots, Cochran's Q statistic²⁹, I² statistic³⁰ and Egger tests³¹ are further discussed in
the eMethods.

181 Moderator analyses were conducted to assess the influence of clinical stage (i.e., CHR-P, EP, 182 and CP), age (i.e., mean age across the study), sex (i.e., proportion of males in the study), 183 antipsychotic medications (i.e., proportion of each study sample taking antipsychotics) and 184 positive and negative symptoms severity on sleep parameters using linear mixed effect meta-185 analysis models. For each sleep parameter, we regressed the differences between patient and 186 control groups from the available sample on each moderator variable. For prevalence, we 187 regressed the log odds of sleep disturbances on each moderator variable. To further investigate 188 the interactions of age, sex, and medication with staging, we applied linear mixed effect models 189 regressing sleep parameter differences on age, sex, medication, and positive and negative 190 symptom severity moderator variables for each stage separately. A threshold of P < 0.017 was 191 used to establish statistical significance after correcting for multiple comparison for three 192 explanatory variables (i.e., sex, age, and proportion medicated) using Bonferroni correction. We 193 also examined the effects of psychotic symptoms (i.e., Positive and Negative Syndrome Scale 194 [PANSS]), and a Bonferroni corrected P threshold of 0.025 was used to establish statistical 195 significance.

196 Results

197 The initial search yielded 7418 records (Figure 1). After removing duplicates, 4863 publications

198 were screened, resulting in 236 studies considered for full-text review. Twelve additional articles

199 were identified through reference checking. After a full-text review, 59 articles were included,

with 21 studies assessing sleep disturbance prevalence in 5135 patients (eTable 1) and 39 studies

201 measuring sleep alterations subjectively (e.g., sleep quality) and/or objectively (e.g., sleep

architecture and sleep oscillatory measures) in 1575 patients and 977 controls (eTable 2).

203 Prevalence of sleep disturbances and insomnia

The pooled (i.e., combined CHR-P, EP and CP) prevalence of sleep disturbances was 50% across clinical stages (95% CI 40% to 61%, Q = 611.28, df = 20, Figure 2A). Stage-specific analyses yielded a sleep disturbance prevalence of 54% in CHR-P (95% CI 40% to 67%, Q = 13.66, df = 3), 68% in EP (95% CI 32% to 90%, Q = 21.2, df = 3), and 44% in CP (95% CI 32% to 57%, Q = 432.73, df = 12, Figure 2A); eFigure 1 shows forest plots of individual studies. Furthermore, prevalence of insomnia as the primary sleep disturbance was 34% (95% CI 24% to 45%) of pooled cases, 48% (95% CI 37% to 59%) of EP, and 27% (95% CI 20% to 35%) of CP (Figure

211 2B, see eFigure 2 for individual studies forest plot). Moderator analysis yielded no sleep

212 disturbance or insomnia differences between clinical stages (eTable 3).

213 Standardized mean differences in sleep quality

Sleep quality was assessed comparing total PSQI scores between clinical and control groups. Results indicated a significant SMD in pooled cases versus controls (SMD [95% CI] = 1.00 [0.70, 1.30], P < 0.001). Each clinical group showed poorer sleep quality compared to controls (CHR-P vs control: SMD [95% CI] = 1.25 [0.83, 1.67], P < 0.001; EP vs control: SMD [95% CI] = 1.17 [0.33, 2.01], P = 0.006; CP vs control: SMD [95% CI] = 0.65 [0.4, 0.89], P < 0.001;

- Figure 3, see eFigure 3 for forest plot of individual studies). Moderator analysis revealed no
 PSQI scores differences between different clinical stages (eTable 3).
- 221 Standardized mean differences in sleep architecture

222 Pooled cases had higher effect sizes for sleep onset latency (SMD = 0.96 [0.62, 1.30], P < 223 0.001), wake after sleep onset (SMD = 0.50 [0.29, 0.71], P < 0.001), number of arousals (SMD 224 = 0.45 [0.07, 0.83], P = 0.019), Stage 1 NREM sleep (SMD = 0.23 [0.06, 0.40], P = 0.008), and 225 REM density (SMD = 0.37 [0.14, 0.60], P = 0.002) vs. controls. Conversely, effect sizes were 226 lower in pooled cases vs control groups for sleep efficiency (SMD = -0.75 [-0.98, -0.52], P < 227 0.001) and slow wave sleep (SMD = -0.24 [-0.44, -0.03], P = 0.023). Furthermore, total sleep 228 time, Stage 2 sleep, and REM latency did not differ between groups (Figure 4; eFigures 4-13 229 contain forest plots of studies for each sleep architecture variable).

230 Stage-specific case-control comparisons revealed no sleep architecture differences in CHR-P vs. controls. EP had higher sleep onset latency (SMD = 0.72 [0.52, 0.92], P < 0.001), wake after 231 232 sleep onset (SMD = 0.62 [0.34, 0.89], P < 0.001), and Stage 1 (SMD = 0.34 [0.15, 0.53], P < 233 0.001), along with lower total sleep time (SMD = -0.56 [-0.99, -0.12], P = 0.012) and sleep efficiency (SMD = -0.90 [-1.20, -0.60], P < 0.001) compared to controls. CP showed higher 234 235 sleep onset latency (SMD = 1.36 [0.66, 2.05], P < 0.001) and wake after sleep onset (SMD = 236 0.51 [0.09, 0.93], P = 0.018), combined with lower sleep efficiency (SMD = -0.73 [-1.14, -0.33], 237 P < 0.001) vs. controls. CP also showed more arousals (SMD = 0.81 [0.30, 1.32], P = 0.002) and 238 REM density (SMD = 0.48 [0.19, 0.77], P = 0.001) compared to controls. Moderator analysis 239 revealed more frequent arousals in CP compared to CHR-P (z = 2.24, p = 0.02, eTable 3).

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- 242 Standardized mean differences in spindle and slow wave parameters
- 243 Pooled cases showed lower spindle density (SMD = -1.06 [-1.50, -0.63], P < 0.001), spindle
- amplitude (SMD = -1.08 [-1.33, -0.82], P < 0.001), and spindle duration (SMD = -1.21 [-1.69, -1.21], -1.69, -1.21
- 245 0.73], P < 0.001, Figure 5) compared to controls. Stage-specific case-control comparisons
- revealed that spindle parameters were lower in both EC and CP relative to controls (Figure 5, see

eFigures 14-16 for forest plots of individual studies). Furthermore, moderator analysis showed

no differences between EP and CP in spindle density or amplitude (eTable 3) but lower spindle

- 249 duration in CP compared to EP (z = -3.91, p < 0.001).
- Finally, slow wave density was not altered in patient groups relative to controls (Figure 5;
 eFigure 17 contains a forest plot of individual studies for slow wave density).
- Supplementary Materials contain results for meta-regressions accounting for medication, age,
 sex, and positive and negative symptoms (eResults and eTables 4-8), study heterogeneity and
 publication bias (eResults, eFigure 18), study quality appraisal (eTables 9-10) and the PRISMA
 2020²⁵ checklist (eTable 11).

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258 Discussion

This meta-analysis investigated sleep abnormalities across clinical stages of psychosis and identified both uniformly present and stage-specific sleep disruptions.

261 Sleep disturbance prevalence is consistently high throughout psychosis stages

Sleep disturbance prevalence has been commonly found to be higher in psychosis compared to the general population^{7,8}, although prior studies reported a variable incidence (21-100%)⁹⁻¹¹. Here, we established that sleep disturbances were present in 50% of pooled clinical cases, with similar prevalence in different psychosis stages, including at-risk individuals. This suggests that sleep disturbances are not only present throughout the course of psychosis, including before the manifestation of a psychotic episode, but are also consistently high in each psychosis stage, thus representing a critical issue that should be addressed in these individuals.

269 Sleep quality is poor throughout stages of psychosis

270 Case-control comparisons of self-reported sleep quality indicated poorer subjective sleep quality 271 in pooled cases and in each clinical stage. Therefore, in addition to sleep disturbances being 272 common, the intensity of perceived sleep distress is also more severe throughout the course of psychosis, including CHR-P, corroborating prior meta-analyses of sleep quality in CHR-P^{9,19}. 273 Notably, in CHR-P poorer sleep quality leads to worse negative symptoms³³ and contributes to 274 predicting transition to psychosis⁵. Together, these findings expose the need to address subjective 275 276 sleep complaints throughout the course of psychosis, even in the at-risk stage. It would therefore 277 be important for primary care and mental health providers to systematically screen for sleep 278 disturbances and to promote sleep hygiene practices (e.g., abstaining from caffeine, nicotine, and alcohol near bedtime, avoiding napping, and maintaining regular sleep and rise times andexposure to daylight) in prodromal individuals.

281 Shared and distinct sleep architecture alterations are present in EP and CP but not in CHR-P

282 Consistent with prior meta-analyses^{15–18}, case-control comparisons of sleep architecture revealed 283 increased sleep onset latency, wake after sleep onset, number of arousals, Stage 1 sleep and REM 284 density, along with lower sleep efficiency and slow wave sleep in pooled clinical cases.

285 Prior work furthermore suggested the presence of specific sleep alterations in early stages of psychosis^{10,19}. However, comparisons from at-risk to chronic stages had thus far not been 286 287 performed. Here, stage-specific case-control comparisons showed that sleep architecture 288 abnormalities were absent in CHR-P and driven by EP and CP stages. Altered sleep 289 characteristics shared among EP and CP included increased sleep onset latency, increased wake 290 after sleep onset, and reduced sleep efficiency. These findings are consistent with insomnia, as 291 well as other disturbances including circadian phase delay, which is supported by recent studies reporting an association between evening chronotype in at-risk ³⁴ and full-blown psychosis³⁵. 292 293 Together, these results suggest that difficulties in initiating and maintaining sleep first occur in 294 full-blown psychosis and remain relatively stable throughout the course of the disorder, as none 295 of the measures worsened in CP vs. EP.

EP also showed a reduction in total sleep time and a higher percentage of Stage 1 sleep, a pattern that was not observed when comparing other clinical stages to their respective control groups. A plausible interpretation of these findings is that individuals in the early course of psychosis suffer from considerable sleep loss and overall shallower sleep, a pattern that is furthermore corroborated by the higher rates of insomnia in EP found in this study. Sleep disruptions in these individuals are also likely involved in psychotic symptomatology, where psychotic experiences worsen sleep and sleep exacerbates psychotic symptoms^{6,36}. From a treatment perspective, earlycourse patients may therefore benefit from routine insomnia screening and targeted sleep interventions, including cognitive-behavioral therapy for insomnia (CBT-I), which is effective in ameliorating difficulties in initiating and maintaining sleep^{37,38}.

306 CP was the only clinical group with more arousals and increased REM density compared to 307 controls. Higher REM density has been associated with increased suicidality in psychotic patients³⁹, and pharmacological reviews indicated that antipsychotic medications can enhance 308 REM density, although effects vary between antipsychotic compounds^{11,40}. Weight gain is a 309 frequent side effect of long-term antipsychotic treatment⁴¹ and has been associated with sleep 310 311 apnea in schizophrenia⁴². Brief awakenings can help restore airflow in such conditions, which 312 may account for the increased frequency of arousals in CP. Moderator analyses further indicated 313 that CP had more arousals compared to CHR-P and that the number of arousals was significantly 314 affected by medication (p=0.001), above and beyond disease effects (z=-3.01 for medication vs. 315 z=2.37 for pooled cases vs. controls). Altogether, these findings indicate that the effects of 316 antipsychotic medications on sleep should be closely monitored, especially in CP, and proper 317 medication adjustments (e.g., decrease medication doses, switch to a different compound) should 318 be considered based on their impact on these sleep patterns.

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322 Meta-analyses of sleep oscillations revealed no alteration in slow wave parameters in clinical 323 cases vs. controls. In contrast, decreased spindle density, spindle amplitude, and duration were 324 observed in pooled cases vs. controls. Stage-specific analyses further indicated that these deficits 325 were present in both EP and CP and yielded some of the largest effect sizes in case-control 326 comparisons (z=-4.93 to -8.31). Of note, spindle measures could not be assessed in CHR-P, as only one study reported spindle measures in this group⁴³. Moderator analyses further indicated 327 328 that CP patients showed a more pronounced reduction of spindle duration compared to EP. 329 Worsening of spindle deficits in chronic stages of psychosis were also reported by a recent metaanalysis on sleep spindles²³, although the clinical groups (schizophrenia, first-episode psychosis, 330 331 and familial risk) and spindle measure (spindle density) only partially overlapped with our study. 332 Furthermore, our moderator analyses revealed considerably larger effect sizes in spindle 333 measures for case-control comparisons (z=-4.93 to -8.31) relative to the effect sizes for the 334 proportion medicated (z=-1.14 to -2.13), and prior studies have consistently shown an absence of correlation between antipsychotic medication and spindle deficits in chronic patients^{44,45}. 335 336 Together, these findings indicate that spindle deficits are unlikely to be related to antipsychotic 337 medications and may represent a neurophysiological biomarker that could be used to monitor the 338 course of psychotic disorders. Furthermore, given increasing evidence for an association between spindle deficits and clinical and cognitive dysfunction in individuals with psychosis^{24,46}, spindles 339 may represent a promising target for novel treatment interventions. In this context, non-invasive 340 brain stimulation has shown promise to restore sleep oscillations, including spindles⁴⁷. 341

344 The current meta-analysis presents some limitations. First, while included studies were selected 345 based on comparable sleep assessment tools, substantial variability in across-study methodology 346 remained. This was most pronounced in sleep disturbance prevalence studies, as sleep disorders 347 were assessed with established diagnostic tools (e.g., DSM criteria for insomnia) in only one study⁴⁸. Similarly, across-study methods employed to measure sleep oscillations varied 348 349 considerably (e.g., manual vs. automated spindle detection, 2-256 electrodes). Notwithstanding 350 this methodological variability, we reported robust, consistent findings, especially regarding 351 spindle deficits in clinical vs. control groups. Second, a few of the included studies were rated as 352 "poor" (N=2), and several were rated as "weak" (N=26). However, the quality of most of these 353 studies was "good" (N=27) or "excellent" (N=5). Third, some analyses had limited statistical 354 power. Specifically, pooled sample sizes of clinical groups included larger samples in CHR-P 355 and CP individuals in prevalence studies, with relatively few prevalence studies in EP. 356 Conversely, for sleep quality and architecture studies, CHR-P sample sizes were smaller 357 compared to EP and CP, indicating that objectively measured sleep is understudied in at-risk 358 populations. The same applied to sleep spindles, which were reported in only one study in CHR- P^{43} . Fourth, due to insufficient data availability, spindles were not stratified into fast and slow 359 360 spindles, although some evidence suggests that distinct alterations in these two types of spindles may occur in psychosis⁴⁹ (see eDiscussion). Similarly, while acute psychosis is likely associated 361 with specific sleep alterations^{50,51}, insufficient data was available to incorporate this factor in the 362 363 current meta-analysis. Fifth, the sleep assessments presented here were based on cross-sectional 364 data rather than on longitudinal evaluations. Nonetheless, this meta-analysis represents the most

365 comprehensive effort to date to delineate sleep abnormalities along the course of psychosis, from366 at-risk to chronic stages.

367 *Conclusion*

This study demonstrates that sleep disturbances are highly prevalent throughout the course of psychosis and that different stages of psychosis show both shared and distinct abnormalities in sleep quality, sleep architecture, and sleep spindle parameters.

371 Altogether, these findings indicate several prospective research directions. To begin with, future 372 studies should use standardized, validated tools to report the prevalence of well-established sleep 373 disorders, as these are common in psychosis but have been rarely assessed in specific clinical 374 stages of psychosis. Moreover, longitudinal sleep studies following at-risk populations through 375 illness stages are necessary to further characterize the interplay between sleep abnormalities and 376 psychosis. To achieve this, research efforts should move beyond conventionally assessed sleep 377 measures and evaluate different sleep patterns using emerging mobile technologies to assess sleep in the home environment³. Future work is also needed to further delineate sleep alterations 378 379 specific to acute and remitted psychosis, as well as the impact of psychotic symptoms severity. 380 Additionally, future studies should better understand the role of antipsychotic medications and 381 different medication types throughout different stages of psychosis. Finally, given the pervasive 382 spindle alterations in EP and CP, an important future direction involves examining spindle properties from CHR-P⁵² to CP stages to determine whether spindle alterations may represent 383 384 risk/susceptibility, monitoring, and/or prognostic biomarkers for psychosis. In doing so, studies 385 should differentiate between fast and slow spindles to accurately delineate psychosis-related

386 sleep alterations. Findings from these studies will help establish sleep as a core clinical target and387 research domain from prodromal to early and chronic stages of psychosis.

388 Author Contributions

- 389 Concept and design: Fabio Ferrarelli, Ahmad Mayeli, and Joëlle Bagautdinova.
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- 391 Drafting of the manuscript: Joëlle Bagautdinova, Ahmad Mayeli, James D. Wilson, Francesco
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- 409 the study and takes responsibility for the integrity of the data and the accuracy of the data

410 analysis.

411 Conflicts of interest

- 412 The authors have no conflicts of interest to declare that are relevant to the content of this article.
- 413
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573 Figure Legends

Figure 1. PRISMA workflow of study selection. *Of note, one study³² was included in both the
 prevalence and sleep architecture analyses.

577 **Figure 2.** Forest plots of A) Prevalence of sleep disturbance and B) Prevalence of insomnia in 578 the pooled clinical groups and each psychosis subgroup. Logit transformation was applied for 579 analysis and the final pooled logit was back transformed to proportions for ease of interpretation 580 of the forest plots.

581

Figure 3. Summary of standardized mean differences in sleep quality as measured by total PSQI
 in the pooled clinical groups and each psychosis subgroup.

584

Figure 4. Summary of standardized mean differences for sleep architecture parameters in pooled clinical groups and in each clinical subgroup. Significant (p < 0.05) effect sizes between two subgroups are marked with an asterisk.

588

589 **Figure 5.** Summary of standardized mean differences for sleep spindle parameters and slow-590 wave density in pooled cases and in each clinical subgroup. Significant (p < 0.05) effect sizes 591 between two subgroups are marked with an asterisk.





B)

Variable	Number of: Patients	
Prevalence		
Insomnia, Pooled Clinical Groups	2740	▶ ──■ ───→
Insomnia, EP	73	⊢ ,
Insomnia, CP	2656	⊢ ⊒ 1
		0.00 0.25 0.50 0.75
		Proportion [05% CI]

Proportion [95% CI]

Proportion [95% CI]

0.50 [0.40, 0.61] 0.54 [0.40, 0.67] 0.68 [0.32, 0.90] 0.44 [0.32, 0.57]

1.00

Proportion [95% CI]

- 0.34 [0.24, 0.45]
- 0.48 [0.37, 0.59]
- 0.27 [0.20, 0.35]

1.00



Variable	Num Patients	ber of: Controls			
Sleen architecture				P Value	SMD [95% CI]
Sleep alcintecture				, ruido	
Total sleep time, Pooled Clinical Groups	631	612		0.206	-0.19 [-0.49, 0.10]
Total sleep time, CHR-P	87	93		0.791	-0.04 [-0.34, 0.26]
Total sleep time, EP	224	219		0.012	-0.56 [-0.99, -0.12]
Total sleep time, CP	320	300		0.8	0.06 [-0.42, 0.54]
Sleep Onset Latency, Pooled Clinical Groups	604	591		<0.001	0.96 [0.62, 1.30]
Sleep Onset Latency, CHR-P	51	62		0.165	0.27 [-0.11, 0.65]
Sleep Onset Latency, EP	231	228		<0.001	0.72[0.52, 0.92]
Sleep Onset Latency, CP	322	301	'⊢'►	<0.001	1.36 [0.66, 2.05]
Sleep Efficiency, Pooled Clinical Groups	608	607	⊢	<0.001	-0.75 [-0.98, -0.52]
Sleep Efficiency, CHR-P	65	73	· · · · · · · · · · · · · · · · · · ·	0.085	-0.30 [-0.65, 0.04]
Sleep Efficiency, EP	239	250		<0.001	-0.90 [-1.20, -0.60]
Sleep Efficiency, CP	304	284	` ⊢=	< 0.001	-0.73 [-1.14, -0.33]
Wake After Sleep Onset, Pooled Clinical Grou	ps 433	431	⊢	<0.001	0.50 [0.29, 0.71]
Wake After Sleep Onset, CHR-P	65	73		0.189	0.23 [-0.11, 0.57]
Wake After Sleep Onset, EP	172	179	' ⊢∎ _	<0.001	0.62 [0.34, 0.89]
Wake After Sleep Onset, CP	196	179	⊢	0.018	0.51 [0.09, 0.93]
Number of arousals, Pooled Clinical Groups	129	124	⊢	0.019	0.45 [0.07, 0.83]
Number of arousals, CHR-P	29	40		0.708	-0.09 [-0.59, 0.40]
Number of arousals, EP	65	54	*	0.09	0.63 [-0.10, 1.35]
Number of arousals, CP	35	30	L '	0.002	0.81 [0.30, 1.32]
Stage 1 Sleep %, Pooled Clinical Groups	536	534	-■-	0.008	0.23 [0.06, 0.40]
Stage 1 Sleep %, CHR-P	51	62		0.405	-0.18 [-0.62, 0.25]
Stage 1 Sleep %, EP	246	253		<0.001	0.34 [0.15, 0.53]
Stage 1 Sleep %, CP	239	219		0.146	0.25 [-0.09, 0.59]
Stage 2 Sleep %, Pooled Clinical Groups	553	551	. F	0.945	0.01 [-0.24, 0.26]
Stage 2 Sleep %, CHR-P	51	62		0.605	-0.10 [-0.48, 0.28]
Stage 2 Sleep %, EP	246	253		0.463	-0.13 [-0.47, 0.22]
Stage 2 Sleep %, CP	256	236		0.395	0.20 [-0.26, 0.66]
Slow Wave Stage %, Pooled Clinical Groups	553	551	⊢∎_	0.023	-0.24 [-0.44, -0.03]
Slow Wave Stage %, CHR-P	51	62		0.547	0.12 [-0.26, 0.50]
Slow Wave Stage %, EP	246	253		0.085	-0.24 [-0.52, 0.03]
Slow Wave Stage %, CP	256	236	⊢	0.109	-0.33 [-0.74, 0.07]
REM Latency, Pooled Clinical Groups	418	416	, F	0.439	-0.09 [-0.32, 0.14]
REM Latency, CHR-P	29	42		0.493	0.31 [-0.58, 1.21]
REM Latency, EP	198	199		0.09	-0.26 [-0.55, 0.04]
REM Latency, CP	191	175		0.799	0.05 [-0.35, 0.45]
REM Density, Pooled Clinical Groups	154	144	·	0.002	0.37 [0.14, 0.60]
REM Density, EP	57	52		0.374	0.17 [-0.21, 0.55]
REM Density, CP	97	92		0.001	0.46[0.19, 0.77]
		Γ			
		-2	-1 0 1 2		
		2	Greater in controls Greater'in patients SMD [95% CI]		

