

the theoretical relevance of these variables to depression the final model achieved an average accuracy of 71% (with individual trials ranging from 64.5% to 77.1%). Key predictors included exploratory behaviors and heart-rate variability during both exploration and cognitive tasks.

Conclusions: These results are comparable, however remain below the levels of accuracy achieved based on fMRI and DTI data alone (around 80%). Nonetheless, the EXPERIENCE system, slated for refinement beyond this pilot phase, shows potential in integrating multimodal data for evaluating affective disorder symptoms, aiming for a more objective screening and diagnostic approach at a lower cost.

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SP0031

Effectiveness and usability of an e-health system on depression among patients with somatic disorders

T. Vitcheva^{1*}, N.G. Petros, G. Hadlaczky and V. Carli

¹Karolinska Institutet, Stockholm, Sweden

*Corresponding author.

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Abstract

Introduction: An increase in the prevalence of depressive symptoms can be seen in patients with severe somatic conditions, with a reduction in quality of life, an increase in sleep disturbances and an increased risk of suicide as some of the most serious consequences. However, few evidence-based interventions have been developed with the aim of reducing this comorbidity. The NEVERMIND system aims to address this issue by collecting psychometric and biomedical data via a smart shirt and a mobile app, which are used to predict patients' depressive symptoms. Patients are then directed to personalised lifestyle behavioural advice, mindfulness-based therapy, and cognitive behavioural therapy.

Objectives: The primary objective was to evaluate the effectiveness of the NEVERMIND system in reducing depressive symptoms in patients with somatic conditions compared to treatment as usual. Secondary objectives included the system's effectiveness in preventing depressive symptoms, sustaining the effects at 24 weeks post-baseline, and reducing suicide ideation. Besides these, the usability, acceptability, and satisfaction of the system were examined in patients with breast or prostate cancer.

Methods: For this pragmatic randomised controlled trial, 425 patients diagnosed with myocardial infarction, breast or prostate cancer, kidney failure, or lower limb amputation were recruited from hospitals in Turin, Pisa and Lisbon. Data collection occurred at baseline, 12 weeks, and 24 weeks, with the primary outcome being depressive symptoms at week 12, measured by the Beck Depression Inventory II. Regarding the usability, acceptability and patient satisfaction, data from 288 patients was used.

Results: The intervention group included 213 and the control group 212 patients, with the sample's mean age being 59.41 (SD=10.70). Patients who used the system reported having statistically significant lower depressive symptoms at 12 weeks (mean difference=-3.05, $p=0.004$; 95%CI -5.12 to -0.99) compared to controls, with a clinically relevant effect size (Cohen's $d=0.41$). Furthermore, significant reductions were found for suicide ideation (mean difference=-0.61, $p=0.020$; 95%CI -1.13 to -0.10) and incidence of depressive symptoms at week 12 (OR=0.43, $p=0.019$; 95%CI 0.22 to 0.87). The decrease in depressive symptoms was sustained at week 24 (mean difference=-1.34, $p=0.015$; 95%CI -2.41 to -0.26). The system was found to have good usability, with women rating the system more favourably than men and valuing its emotional support, while men used the system more frequently than women and valued the self-awareness that the system encouraged.

Conclusions: The NEVERMIND system was shown to be superior to standard care in reducing and preventing depressive symptoms among the studied sample. A new project will be launched in the near future to continue the examination of the system's effectiveness.

Disclosure of Interest: None Declared

SP0032

Inflammatory based psychotic symptoms: when psychosis means encephalitis

M. Rojnic Kuzman

Zagreb University Hospital Centre and the Zagreb School of Medicine, Zagreb, Croatia

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Abstract: Schizophrenia, as one of the most common disorders from the psychotic spectrum is most commonly detected in the phase of first psychosis and may pose a diagnostic challenge, as commonly comprise a heterogeneous group of schizophrenias, with distinct clinical presentations. If it detected in its prodromal phase without clearly developed psychotic symptoms, the diagnosis is even more unreliable, as the transition to full blown psychosis in the next two years happens in 15-40% of more, depending probably on a variety of cumulative environmental risk factors (including childhood trauma, the use of high-potency cannabis, urbanicity, season of birth). Moreover, the first episode psychosis may underlie for example the first manic episode, brief intermittent psychotic symptoms in persons with borderline personality disorders, acute reaction to trauma, the use of cannabis and psychostimulants and different organic causes, such as endocrinologic disorders and autoimmune encephalitis. Therefore, in everyday clinical practice, the diagnosis of first episode psychosis always requires an assessment of possible causes of psychosis, and also factors that may influence prognosis and treatment. Usual assessment include detailed anamnestic and heteroanamnestic data, physical examination, standard blood laboratory findings, drugs in urine/ blood, EEG and CT/MR scan. The absence of typical risk factors for schizophrenia, as well as the absence of premorbid symptoms and developmental course typical for schizophrenia, abrupt course of psychotic symptoms, symptoms such as disorientation,

catatonia, speech disturbances, alteration of consciousness, neurologic signs, autonomic dysfunction and laboratory aberrations may be especially indicative for organic cause and possibly encephalitis and require further confirmation with the analysis of cerebrospinal liquor with antineuronal antibodies.

Disclosure of Interest: None Declared

SP0033

Childhood trauma as a predictor of social cognition disturbances across psychosis spectrum: Data from the PREGAP Study

C. Aymerich^{1,2,3,4*} and A. Catalan^{1,2,3,4}

¹Psychiatry, Basurto University Hospital; ²Psychiatry, Biobizkaia Health Research Institute, Bilbao; ³Neurosciences, Basque Country University (UPV/EHU), Leioa and ⁴GCV22/1/SAM, CIBERSAM, Madrid, Spain

*Corresponding author.

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Abstract

Introduction: Childhood trauma is a severe form of stress that has been strongly related to both the appearance of a psychotic disorder and the existence of social cognition disturbances. We hereby hypothesize childhood trauma might be a transdiagnostic marker of social cognition disturbances across the psychosis spectrum, regardless of the main diagnosis.

Objectives: To investigate the effect of different forms of childhood trauma in social cognition impairments in first-episode psychosis, at-risk mental states for psychosis and healthy controls.

Methods: Using cross-sectional data, we will examine the relationship between different kinds of childhood trauma (measured with the Childhood Trauma Questionnaire, CTQ) and several social cognition domains, including facial emotion recognition, theory of mind (assessed using the Movie Assessment for Social Cognition, MASC, The Hinting Task, and the Faux-Pas Questionnaire). Intra and inter-group differences be studied for three study groups, including patients with first-episode psychosis (n=60), subjects with at-risk mental states for psychosis (n=60), and healthy controls (n=60).

Disclosure of Interest: None Declared

SP0034

Childhood trauma as a transdiagnostic risk factor: clinical implications and preventive interventions

R. J. Bou Khalil

Psychiatry, Saint-Joseph University, Beirut, Lebanon

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Abstract: Abstract: This presentation seeks to explore the interplay between various types of psychological traumas and their potential correlation with the development of distinct types and severities of

eating disorders. Emphasis will be placed on elucidating the underlying biological underpinnings and psychological and developmental factors that contribute to the manifestation of diverse eating disorder phenotypes in individuals who have experienced childhood maltreatment.

Drawing upon existing research and novel insights, I will present some data from studies investigating the notion that the observed variations in eating disorder presentations may be linked especially to environmental influences. Contrary to the conventional focus on genetic determinants, our findings suggest that the differential ecophenotypic expression of eating disorders may not solely be attributed to DNA variants but rather to the complex interplay between genetic predispositions and environmental contexts.

In particular, I will expose the concept of an ecophenotype characteristic of eating disorders associated with childhood maltreatment, positing that the unique ecological context in which an individual is raised significantly influences the trajectory and severity of their eating disorder. This exploration extends beyond a mere examination of genetic markers, shedding light on the environmental and ecosystemic factors that shape the development of an individual's relationship with food and body image.

Disclosure of Interest: None Declared

SP0035

How is trauma a transdiagnostic risk factor? A biopsychosocial model of risk and protective mechanisms following childhood trauma

H. Kerbage^{1,2*} and D. Purper-Ouakil^{1,2}

¹University Hospital Center of Montpellier (CHU de Montpellier), Montpellier and ²Center for Epidemiology and Population Health (CESP), Paris, France

*Corresponding author.

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Abstract: Traumatic exposure is a common global problem across nations. It is currently well established that childhood trauma is associated with increased risk for psychopathology transdiagnostically, with children having experienced trauma being twice as likely to develop a mental health condition compared to those who have never experienced trauma. According to population-based studies, this heightened risk for the emergence of mental health disorders persists throughout adolescence and adulthood. The risk for psychopathology seems to be most marked in children exposed to interpersonal violence (child emotional and physical abuse, neglect, sexual violence). In this presentation, we will summarize the results of an increasing number of published studies that have examined the mechanisms underlying vulnerability to psychopathology following childhood trauma and protective factors that buffer this risk. Specifically, we will highlight the role of emotion dysregulation and interpersonal difficulties, related to disrupted threat processing following trauma exposure, in mediating the impact of trauma on internalizing and externalizing symptoms. Research studies have also identified protective factors across the lifespan that might mitigate these outcomes, including social support and emotional skills building. Based on this review, we will suggest a conceptual transdiagnostic and biopsychosocial model of risk and resilience, which can provide opportunities and targets for early interventions and treatment, at the