

## Joint Symposium

### JS0001

#### Dysregulated metabolic sensing of appetite in anorexia nervosa: implications of LEAP-2 regulation

C. Tezenas Du Montcel<sup>1,2\*</sup>

<sup>1</sup>Genetic and epigenetic vulnerabilities to addictive and psychiatric disorder, Inserm UMR 1266 and <sup>2</sup>Cliniques des Maladies mentales et de l'Encéphale, GHU Paris Psychiatrie et Neurosciences, Paris, France

\*Corresponding author.

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**Abstract:** Growing interests on the role of metabolic sensors in anorexia nervosa led to implicate metabolic sensing as consequences of anorectic sensing but also in the perpetuation of the disorder. Ghrelin is an orexigenic peptide secreted by the fundic cells of the stomach in situation of fasting and known to initiate food intake through its activity on hypothalamic and motivation aspect of food intake. A body of evidence previously showed that patients suffering from anorexia nervosa display high plasma levels of ghrelin correlated with the nutritional status but this orexigenic signal do not seem to modify restrictive behavior. LEAP-2 (Liver Expressed Antimicrobial Peptide 2) is a recently discovered endogenous ghrelin antagonist, increased during overnutrition and that decreases food intake in humans and animals.

We explored changes of ghrelin and LEAP-2 in a longitudinal cohort of 30 patients suffering from anorexia nervosa during a 4 months refeeding program. We show abnormal regulation of LEAP-2 in patients with higher levels in acute stages that decrease with refeeding. This abnormal regulation was associated with early relapse in patients. This abnormal regulation could counteract with the orexigenic signal of ghrelin in patients.

We discuss these results in light with recent evidence on the consequences of LEAP-2 increase of food intake and hedonic feeding relevant in understanding anorexia nervosa.

**Disclosure of Interest:** None Declared

### JS0002

#### Potential Neurobiological and clinical markers in Extreme Weight Conditions: from Anorexia to Obesity

F. Fernandez-Aranda<sup>1,2,3\*</sup>, S. Jimenez- Murcia<sup>1,2,3</sup> and Psychoneurobiology of Eating and Addictive Behaviours Group, Neuroscience Program, IDIBELL

<sup>1</sup>Clinical Psychology, University Hospital of Bellvitge-IDIBELL;

<sup>2</sup>CIBERObn, Instituto Salud Carlos III and <sup>3</sup>Clinical Sciences, University of Barcelona, Barcelona, Spain

\*Corresponding author.

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**Abstract:** Extreme eating and weight conditions (EWC) are a construct that emerges as a dimensional and theoretical model that identifies individuals who exhibit inappropriate eating behaviours and abrupt weight fluctuations. According to this spectrum of

EWC, one extreme can be represented by individuals with anorexia nervosa (AN), characterised by excessive food restriction and an extremely low body mass index (BMI), whereas the other end of this continuum is represented by individuals with obesity (OB), characterised by a BMI above 30. In addition to AN and OB, some eating disorders (EDs), namely bulimia nervosa and binge eating disorder, are also part of this continuum, given the high risk of falling into one of the extremes, especially that of higher BMI. Studies have described similar changes at the psychological and neurobiological levels associated with their abnormal eating patterns, delineating vulnerability pathways related to the neurobiological basis.

Based on previous literature, individuals suffering from EWC would show dysfunctional brain activity in regions associated with emotional reward processing and cognitive control compared to healthy controls (HC). Similarly, neuroendocrine alterations in EWC are expected to influence clinical symptomatology. It will also be discussed how impairments in executive function and differential brain activity observed in individuals with EWC may negatively impact their clinical course and treatment outcome.

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### JS0003

#### (Ir)reversibility of structural and functional brain alterations in severe anorexia

L.-K. Kaufmann

Radboud University, Donders Institute for Brain, Cognition and Behaviour, Nijmegen, Netherlands

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**Abstract:** Anorexia nervosa is characterized by profound structural and functional brain alterations, particularly during the phase of acute underweight. Understanding the reversibility of these changes upon weight normalization is an important question in the pursuit of recovery and relapse prevention. This talk shares findings from recent neuroimaging studies, focussing on the dynamic processes of brain recovery observed during and after inpatient treatment in individuals with severe anorexia nervosa.

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