Evaluation of plasma cytokines in patients with cocaine use disorders in abstinence identifies transforming growth factor alpha (TGFα) as a potential biomarker of consumption and dual diagnosis

Rosa Maza-Quiroga1,*, Nuria García-Marchena1,*, Pablo Romero-Sanchiz1,*, Vicente Barrios3, María Pedraz1, Antonia Serrano1, Raquel Nogueira-Arjona1, Juan Jesus Ruiz3, Maribel Soria3, Rafael Campos3, Julie Ann Chowen2, Jesus Argente2, Marta Torrens4, Meritxell López-Gallardo5, Eva María Marco6, Fernando Rodríguez de Fonseca1, Francisco Javier Pavón1 and Pedro Araos6,*

1 Hospital Regional Universitario de Málaga, Unidad de Gestión Clínica de Salud Mental, Instituto de Investigación Biomédica de Málaga (IBIMA), Málaga, Spain
2 Department of Endocrinology, Hospital Universitario Niño Jesús, Madrid, Spain
3 Diputación de Málaga, Centro Provincial de Drogedependencias, Málaga, Spain
4 Institut de Neuropsiquiatria i Addiccions (INAD) del Parc de Salut Mar, Barcelona, Spain
5 Department of Physiology Faculty of Medicine, Universidad Complutense de Madrid, Madrid, Spain
6 Department of Physiology II Faculty of Biology, Universidad Complutense de Madrid, Madrid, Spain
* These authors contributed equally to this work.

ABSTRACT

Background. Cocaine use disorder (CUD) is a complex health condition, especially when it is accompanied by comorbid psychiatric disorders (dual diagnosis). Dual diagnosis is associated with difficulties in the stratification and treatment of patients. One of the major challenges in clinical practice of addiction psychiatry is the lack of objective biological markers that indicate the degree of consumption, severity of addiction, level of toxicity and response to treatment in patients with CUD. These potential biomarkers would be fundamental players in the diagnosis, stratification, prognosis and therapeutic orientation in addiction. Due to growing evidence of the involvement of the immune system in addiction and psychiatric disorders, we tested the hypothesis that patients with CUD in abstinence might have altered circulating levels of signaling proteins related to systemic inflammation.

Methods. The study was designed as a cross-sectional study of CUD treatment-seeking patients. These patients were recruited from outpatient programs in the province of Malaga (Spain). The study was performed with a total of 160 white Caucasian subjects, who were divided into the following groups: patients diagnosed with CUD in abstinence (N = 79, cocaine group) and matched control subjects (N = 81, control group). Participants were clinically evaluated with the diagnostic interview PRISM according to the DSM-IV-TR, and blood samples were collected for the determination of chemokine C-C motif ligand 11 (CCL11, eotaxin-1), interferon gamma (IFNγ), interleukin-4 (IL-4), interleukin-8 (IL-8), interleukin-17α (IL-17α), macrophage inflammatory...
protein 1α (MIP-1α) and transforming growth factor α (TGFα) levels in the plasma. Clinical and biochemical data were analyzed in order to find relationships between variables.

**Results.** While 57% of patients with CUD were diagnosed with dual diagnosis, approximately 73% of patients had other substance use disorders. Cocaine patients displayed greater cocaine symptom severity when they were diagnosed with psychiatric comorbidity. Regarding inflammatory factors, we observed significantly lower plasma levels of IL-17α ($p < 0.001$), MIP-1α ($p < 0.001$) and TGFα ($p < 0.05$) in the cocaine group compared with the levels in the control group. Finally, there was a significant primary effect of dual diagnosis on the plasma concentrations of TGFα ($p < 0.05$) in the cocaine group, and these levels were lower in patients with dual diagnoses.

**Discussion.** IL-17α, MIP-1α and TGFα levels are different between the cocaine and control groups, and TGFα levels facilitate the identification of patients with dual diagnosis. Because TGFα reduction is associated with enhanced responses to cocaine in preclinical models, we propose TGFα as a potential biomarker of complex CUD in humans.

**Subjects** Cognitive Disorders, Neurology, Psychiatry and Psychology  
**Keywords** Cocaine use disorders, Cytokines, Dual diagnosis

**INTRODUCTION**

Cocaine use disorder (CUD) constitutes a highly complex health problem involving not only biological changes in the brain but also a variety of social and environmental aspects (Tomasi et al., 2015). The chronic use of cocaine is often accompanied by medical issues, including circulatory diseases and comorbid psychiatric conditions (González-Saiz et al., 2014; Verdejo-Garcia et al., 2015; Chibungu et al., 2016). Dual diagnosis or co-occurring disorders describes the presence of both a mental health and a substance use disorder, and the dual nature of this condition complicates the diagnosis and therapeutic outcomes in CUD because many of the patients do not respond to classical pharmacological approaches (Torrens, Gilchrist & Domingo-Salvany, the PsyCoBarcelona Group, 2011; Álvarez et al., 2013). Moreover, the quality of life of these patients is severely impaired (Chahua et al., 2015).

To improve the therapeutic process in addiction, one of the challenges is to identify biological markers that might aid in objectively determining the degree of consumption, severity of addiction, level of toxicity and response to treatment in patients with CUD (Pavón et al., 2013; Araos et al., 2015). These potential biomarkers would be fundamental players in the diagnosis, stratification, prognosis and therapeutic orientation in addictions. Several attempts have been made to identify these possible biomarkers by exploring accessible signaling compartments (i.e., blood) or functional neuropsychological tests. Among these attempts, recent studies have identified circulating factors involved in immune function as potential biomarkers in CUD and other substance use disorders (Araos et al., 2015; Moreira et al., 2016; García-Marchena et al., 2017a). These and other studies have demonstrated that abused drugs interact with the immune system and alter signaling and gene expression.
involved in the immune response, which these effects contribute to various aspects of addiction (Cui, Shurtleff & Harris, 2014).

More specifically, cocaine abuse results in increased pro-inflammatory signaling throughout the brain (Cearley et al., 2011; Coller & Hutchinson, 2012); however, the mechanism is unknown. These alterations are also found in some plasma circulating chemokines and cytokines (Araos et al., 2015), indicating that alterations in pro- and anti-inflammatory factors may serve as potential biomarkers of an inflammatory state in the central nervous system (CNS). In fact, this systemic inflammation has been recently proposed to be a mechanism that facilitates dopamine signaling in the brain, which contributes to the addiction cycle (Petrulli et al., 2017).

The fact that the immune system modifies brain functions related to addiction and the concurrent participation of reward modulatory systems in psychiatric disorders open the possibility of establishing a link between inflammation, neuropsychiatric diseases and addictive disorders (Stertz, Magalhaes & Kapczinski, 2013). Comorbid mental health and substance use disorders are usually present in more than 50% of cocaine- or alcohol-addicted patients (Vergara-Moragues et al., 2012; Garcia-Marchena et al., 2017b). Convergence of neuroinflammation, stress and addiction in neural substrates of mental diseases could help us to understand this complex association. Stressful experiences may model the immune system, modifying the release and signaling of cytokines involved in inflammation (Slavich, 2016). In addition, several studies indicate that the inflammatory processes could actively influence the CNS and that this would generate changes in the behavior of the subjects (Irwin & Miller, 2007). As an example, neuroinflammation and alterations in neurogenesis are clear substrates of mood disorders (Goldstein et al., 2009), and both processes are clearly affected by abused drugs (Castilla-Ortega et al., 2016; Neupane, 2016).

This research is the continuation of a previous study in patients with CUD (Araos et al., 2015); in the present study, we also analyzed the inflammatory signaling factors in the blood of patients addicted to cocaine to explore the relationships between the expression of circulating agents related to neuroinflammation and neurogenesis (cytokines, chemokines and growth factors) and variables associated with CUD and dual diagnosis. The selected molecules in these two studies were chosen according to an exhaustive bibliographic search.

The inflammatory signals examined in the current study were chemokine C-C motif ligand 11 (CCL11, eotaxin-1), interferon gamma (IFNγ), interleukin-4 (IL-4), interleukin-8 (IL-8), interleukin-17α (IL-17α), macrophage inflammatory protein 1α (MIP-1α) and transforming growth factor α (TGFα).

The main goal of the present study was to evaluate the plasma concentrations of cytokines, chemokines and growth factor in a cohort of CUD patients in abstinence compared to population controls. Variables related to CUD, such as the duration of problematic cocaine use, abstinence length, cocaine addiction severity and dual diagnosis, were also examined in relation to these molecules.
MATERIALS AND METHODS

Study design and recruitment
A cross-sectional study was conducted on patients diagnosed with CUD seeking treatment for cocaine use compared with control subjects. The cocaine patients were recruited from outpatient treatment programs in the province of Malaga (Spain) for a period of 24 months (August 2014 to August 2016). A total of 160 white Caucasian subjects were recruited and divided into the cocaine group and the control group. To be eligible for the study, participants had to be \( \geq 18 \) years to 65 years of age. Exclusion criteria included a personal history of the following: (1) chronic diseases (e.g., cardiovascular, respiratory, renal, hepatic, neurological or endocrinological diseases); (2) cancer; (3) infectious diseases; (4) incapacitating cognitive alterations; and (5) pregnancy for women.

Cocaine group and other dual diagnoses
Seventy-nine abstinent patients were diagnosed with CUD (cocaine abuse and/or dependence). The diagnosis of CUD and other psychiatric disorders were performed with a psychiatric interview (‘Diagnostic and Statistical Manual of Mental Disorders-4th Edition-Text Revision’, DSM-IV-TR), while the abstinence from abused drugs was controlled weekly by urine analysis in the outpatient treatment centers for cocaine addiction. Urine analysis for cocaine, amphetamine, opiates, barbiturates, phencyclidine and cannabis was performed using a V-Twin Drug Testing System (Siemens AG, Erlangen, Germany). These data were used to select patients with CUD who abstained from abused drugs during the last 2 weeks (at least). Subsequently, plasma analyses were conducted to verify cocaine abstinence.

Control group without dual diagnosis
The control group was recruited from a multidisciplinary staff working at the Hospital Regional Universitario de Málaga (Málaga, Spain). Eighty-one healthy and unmedicated participants were matched to the cocaine group for age, sex and body mass index (BMI) in order to provide reference values of circulating inflammatory factors. Subjects with substance use disorders and past or current Axis I and Axis II disorders (DSM-IV-TR) and neurologic disorders were excluded from the control group.

Ethics statement
Written informed consent was obtained from each participant after a complete description of the study and after questions or issues were discussed. The study and protocols for recruitment were approved by the Ethics Committee of the CEI Provincial de Málaga in accordance with the ‘Ethical Principles for Medical Research Involving Human Subjects’ that was adopted in the Declaration of Helsinki by the World Medical Association (64th WMA General Assembly, Fortaleza, Brazil, October 2013). The data were protected by the Recommendation No. R (97) 5 of the Committee of Ministers to Member States on the Protection of Medical Data (1997) and the Spanish data protection act (Ley Orgánica 15/1999 de Protección de Datos, LOPD).
Psychiatric evaluation

All patients with CUD were evaluated according to the DSM-IV-TR criteria using the Spanish version of the ‘Psychiatric Research Interview for Substance and Mental Disorders’ (PRISM 6.0) (Hasin et al., 1996; Torrens et al., 2004). Control subjects were initially evaluated by PRISM to detect substance use disorders and by the Spanish version of ‘Dual Diagnosis Screening Instrument’ (DDSI) to detect psychiatric disorders (DSM-IV-TR Mestre-Pintó et al., 2014). All interviews were conducted by psychologists who had received training certificates for these instruments.

PRISM

The PRISM diagnostic interview was used to assess psychopathological and substance use disorders. PRISM is a semi-structured clinical interview designed to solve the problems of diagnosis in people with consumption of substances and/or alcohol. This interview presents good test-retest reliability, validity and inter-examiner reliability (kappa coefficient oscillates between 0.66 and 1.00) (Morgello et al., 2006). The first module contains questions related to the history of consumption, ranging from the time of the diagnosis of abuse and/or dependence. In addition, PRISM evaluates 20 Axis I disorders and 2 Axis II disorders (borderline and antisocial personality disorders) according to DSM-IV-TR. The diagnoses are performed using two time-frames: current (criteria were met within the past year) and past (criteria were met before the previous 12 months). Consequently, the estimated prevalence of a lifetime diagnosis would include both current and past diagnoses.

In addition, PRISM allows differentiation between primary mental disorders (or independent disorders) and disorders induced by substances in combination with the expected symptoms of the effect of intoxication and abstinence. The criteria established by PRISM for a substance-induced disorder is that the disorder must occur in the context of a pathological consumption of the substance in any of these 2 situations: (a) chronic intoxication (consumption for four or more days a week for a month) and (b) binge (consumption for a period of three continuous days) (Hasin et al., 1996; Torrens et al., 2004).

The cocaine severity symptom was used to determine cocaine trait severity combining the seven dependence criteria and the four abuse criteria (DSM-IV-TR) (Pavón et al., 2013; Araos et al., 2015; Pedraz et al., 2015).

Collection of plasma samples for analysis

Blood samples were extracted in the morning (08:00–10:00 AM) after fasting for 8–12 h and before the psychiatric interview. Venous blood samples were extracted into 10-mL K2 EDTA tubes (BD, Franklin Lakes, NJ, USA) and centrifuged at 2,200 g for 15 min (4 °C) to obtain plasma. Plasma samples were individually assayed by three commercial tests for detecting infectious diseases: HIV, hepatitis B and hepatitis C. Plasma analyses for cocaine metabolite (Benzoylecgonine Specific Direct ELISA Kit purchased from Immunalysis Co., Pomona, CA, USA) were also performed to confirm cocaine abstinence. Plasma samples were stored at −80 °C.
Multiplex immunoassay analysis
For plasma determination, inflammatory proteins were chosen considering those inflammatory factors that were not determined previously in abstinent cocaine patients (see details in Araos et al., 2015). A Bio-Plex Suspension Array System 200 (Bio-Rad Laboratories, Hercules, CA, USA) and ProcartaPlex Immunoassay Kit with magnetic beads and an appropriate Plasma Standard Diluent Kit (eBioscience, Affymetrix, Santa Clara, CA, USA) were used to quantify protein levels in the plasma. This method of analysis is based on the Luminex technology, and a human cytokine 7-plex panel (Mix&Match Panel) was used to simultaneously detect the following analytes: chemokine C-C motif ligand 11 (CCL11, eotaxin-1), interferon gamma (IFNγ), interleukin-4 (IL-4), interleukin-8 (IL-8), interleukin-17α (IL-17α), macrophage inflammatory protein 1α (MIP-1α) and transforming growth factor α (TGFα).

The measurements of these analytes in plasma were performed following the manufacturer’s instructions.

Statistical analysis
All clinical data in Tables 1 and 2 are expressed as the number and percentage of subjects (N (%)) or the mean and SD (mean (SD)). The significance of differences in categorical variables was determined using Fisher’s exact test (Chi-square test). The significance of differences in normal continuous variables or non-normal continuous variables was determined using Student’s t-test and Mann–Whitney U test, respectively. Statistical analysis of protein levels was performed using multiple analysis of covariance (ANCOVA) to indicate the relative effect of explanatory variables and their interactions on the protein expression in the plasma, controlling for additional covariates. Log (10) transformation was used to ensure statistical assumptions for positive skewed distributions. Estimated marginal means (95% confidence intervals [95% CI]) of protein levels were expressed after back-transformation as shown in Table 3 and Figs. 1 and 2.

In this study, we used the Kolmogorov–Smirnov test with Lilliefors correction to analyze normality of data. We used Levene’s test to analyze homoscedasticity of the data.

All statistical analyses were performed using R-commander version 3.3.2 free software and GraphPad Prism version 5.04 software (GraphPad Software, San Diego, CA, USA). A p-value <0.05 was considered statistically significant. The specific statistical analysis used is indicated in the text and in each figure caption.

RESULTS
Socio-demographic characteristics
The description of the socio-demographic variables of the participants is presented in Table 1. A total of 160 subjects met the eligibility criteria for this study and were divided into the cocaine (N = 79) and control (N = 81) groups.

The average age of the participants in the cocaine abstinent group was 35 years old, and the average BMI was 26. We found significant differences between patients with CUD and the controls in the marital status variable (p < 0.05). Cocaine patients had a significantly
Table 1  Baseline socio-demographics and psychiatric characteristics in cocaine and control group.

| Variable                        | Total N = 160 | Cocaine N = 79 | Control N = 81 | p value |
|---------------------------------|---------------|----------------|----------------|---------|
| Age (mean(SD))                  |               | 34.87 (7.18)   | 37.27 (10.97)  | 0.115   |
| BMI (mean(SD))                  |               | 25.88 (4.20)   | 25.32 (3.65)   | 0.370   |
| Sex (N(%))                      |               | 16 (20.25)     | 16 (19.75)     | 1^      |
| Marital status (N(%))           |               | 29 (36.71)     | 26 (47.3)      |         |
| Education (N(%))                |               | 63 (79.75)     | 9 (11.11)      | <0.001  |
| psychiatric treatment (N(%))    |               | 52 (65.82)     | 72 (88.89)     |         |
| Dual diagnosis (N(%))           |               | 34 (43.04)     | 45 (56.96)     |         |
| Other substance use disorders (N(%)) |           | 21 (26.58)     | 58 (73.42)     |         |
| Length of abstinence (mean (SD))|               | 133.4 (114.53) | -              |         |
| Cocaine symptom severity (mean (SD)) |         | 8.04 (2.66)    | -              |         |

Notes.
- BMI, body mass index.
- ^p-value from Student’s-test.
- ^p-value from Wilcoxon-test.
- ^p-value from Chi-square-test.

lower educational level and a higher percentage of psychiatric treatments compared with the control group—(p < 0.001).

A total of 57% of cocaine patients were diagnosed with dual diagnosis, which includes presenting with at least one psychiatric disorder throughout life (i.e., mood, anxiety, psychotic or personality disorders (antisocial and borderline)). In addition, 73% of cocaine patients were diagnosed with comorbid substance use disorders (e.g., alcohol, heroin, cannabis, benzodiazepines, hallucinogens or other stimulants). The length of abstinence from cocaine in these patients was 133 days, and the mean cocaine symptom severity was 8 CUD-criteria at the time of recruitment.

**Plasma levels of inflammatory signaling proteins in subjects with CUD and in controls**

As shown in Fig. 1, the estimated marginal means of eotaxin-1, IFNγ, IL-4, IL-8, IL-17α, MIP-1α and TGFα in the plasma are presented according to the history of cocaine use (the cocaine and control groups). The plasma concentrations of these factors were analyzed by ANCOVA using ‘cocaine use’ as the main factor and controlling for age, sex and BMI.

We found a significant main effect of ‘cocaine use’ on plasma levels of IL-17α –(F_{1,133} = 20.713, p < 0.001), MIP-1α –(F_{1,147} = 26.444, p < 0.001) and TGFα –(F_{1,540} = 5.251, p < 0.05). Thus, cocaine patients showed lower levels of these inflammatory
proteins than the control group. In contrast, we found no differences between both groups in the levels of eotaxin-1, IFNγ, IL-4 or IL-8.

Impact of psychiatric comorbidity on plasma levels of inflammatory signaling proteins in subjects with CUD

Cocaine patients were divided into subgroups according to the diagnosis of psychiatric comorbidity. Additionally, we used two different criteria of classification to explore the effects of psychiatric comorbidity on these inflammatory factors: (a) Diagnosis of comorbid substance use disorders and (b) Dual diagnosis with other mental health disorder(s). A description of these patients is shown in Table 2.

A first classification of the cocaine group was performed according to the diagnosis of other substance use disorders (yes (N = 58); no (N = 21)). The comparison between both subgroups showed that there were no significant differences in age, BMI, sex, psychiatric treatment, age of cocaine initiation or length of abstinence. However, we found significant differences in the cocaine symptom severity –(p < 0.001), and patients with comorbid substance use disorders had higher cocaine severity than patients with no other substance use disorders (8.4 vs. 7.0 criteria). Although patients with substance use disorders had an elevated prevalence of dual diagnosis (64%) compared with patients with no other substance use disorders (38%); however, this difference did not reach statistical significance (p = 0.075).

An additional classification was performed according to the detection of dual diagnosis (yes (N = 45); no (N = 34)). We found no differences in BMI, sex, psychiatric treatment, age of cocaine initiation or length of abstinence. In this case, we observed significant
Table 2  Baseline socio-demographic variables and psychiatric characteristics in substance use disorders and dual diagnosis.

| Variable                                      | Cocaine N = 79 |
|-----------------------------------------------|----------------|
|                                               | Other substance use disorders | Other substance use disorders | p value | Dual diagnosis | Dual diagnosis | p value |
| Participants (N (%))                          | Yes | 58 (73.42) | 21 (26.58) | – | 45 (56.96) | 21 (26.58) | – |
|                                               | No  | 21 (26.58) | 21 (26.58) | – | 34 (43.04) | 34 (43.04) | – |
| Age (mean (SD))                               | Years | 34.72 (7.15) | 35.29 (7.42) | 0.726b | 36.53 (6.65) | 32.68 (7.18) | 0.025b |
| BMI (mean (SD))                               | kg/m² | 25.58 (3.41) | 26.70 (5.89) | 0.300a | 26.58 (4.61) | 24.95 (3.43) | 0.089a |
| Sex (N (%))                                   | Women | 10 (17.24) | 6 (28.58) | 0.340c | 11 (24.44) | 5 (14.71) | 0.433c |
|                                               | Men  | 48 (82.76) | 15 (71.43) | – | 34 (75.56) | 29 (85.29) | – |
| Psychiatric treatment (N(%))                  | No  | 37 (63.79) | 15 (71.43) | 0.716c | 26 (57.78) | 26 (76.47) | 0.135c |
|                                               | Yes | 21 (36.21) | 6 (28.58) | – | 19 (42.22) | 8 (23.53) | – |
| Age of cocaine initiation (mean (SD))         | Years | 26.48 (8.12) | 28.90 (7.47) | 0.236c | 26.42 (7.86) | 28.06 (8.16) | 0.507c |
| Length of abstinence (mean (SD))              | Days | 139.50 (113.20) | 116.30 (119.26) | 0.287c | 133.5 (102.68) | 133.2 (130.19) | 0.334c |
| Cocaine symptom severity (mean (SD))          | (0–11)criteria | 8.41 (2.56) | 7.00 (2.70) | <0.001b | 9.20 (1.65) | 6.50 (2.97) | <0.001b |
| Dual diagnosis (N (%))                        | No  | 21 (36.21) | 13 (61.90) | 0.075c | – | – | – |
|                                               | Yes | 37 (63.79) | 8 (38.10) | – | 8 (17.78) | 13 (39.24) | 0.075c |
| Other substance use disorders (N (%))          | No  | – | – | – | 37 (82.22) | 21 (61.76) | – |
|                                               | Yes | – | – | – | – | – | – |

Notes.
- BMI, body mass index.
- p-value from Student’s-test.
- p-value from Wilcoxon-test.
- p-value from Chi-square-test.

Differences in age (p < 0.05) and in the cocaine symptom severity (p < 0.001). Further, patients with dual diagnosis were older (36.5 vs. 32.7 years) and had a more severe form of CUD (9.2 vs. 6.5 criteria) than patients with no dual diagnosis. Furthermore, these patients with dual diagnosis had an increased prevalence of comorbid substance use disorders (82%) relative to patients with no dual diagnosis (62%), but this difference did not quite reach statistical significance (p = 0.075).

Impact of dual diagnosis on plasma levels of inflammatory signaling proteins in subjects with CUD

Figure 2 shows the estimated marginal means of eotaxin-1, IFNγ, IL-4, IL-8, IL-17α, MIP-1α and TGFα in the plasma of cocaine patients, who were grouped according to the presence of dual diagnosis. The plasma concentrations of these inflammatory markers were analyzed by ANCOVA using ‘dual diagnosis’ as the main factor and controlling for age, sex and BMI.

There was a significant main effect of ‘dual diagnosis’ on the plasma levels of TGFα (F1,190 = 6.812, p < 0.05). Thus, cocaine patients with dual diagnosis showed lower TGFα levels than patients with no dual diagnosis. Consequently, the presence of dual diagnosis
in the cocaine group enhanced a decrease in TGFα levels relative to the levels in the control group.

Impact of cocaine symptom severity on plasma levels of inflammatory signaling proteins in subjects with CUD

Because we observed significant differences in the cocaine symptom severity when cocaine patients were grouped according to psychiatric comorbidity (comorbid substance use disorders and dual diagnosis), we explored the impact of the degree of cocaine severity on these circulating inflammatory proteins. Therefore, the cocaine group was divided into 2 subgroups as follows: mild/moderate ((0–8 criteria) \(N = 38\)) and severe ((9–11 criteria) \(N = 41\)) CUD.
Similar to previous analyses, plasma concentrations of these proteins were analyzed by ANCOVA using ‘cocaine severity’ as the main factor and controlling for age, sex and BMI. However, as shown in Table 3, we found no differences between both subgroups.

DISCUSSION

The search for potential biomarkers might enable a better stratification of patients to establish therapeutic subgroups. These biomarkers might arise from the activity of biological systems involved in the pathogenesis of mental disorders. There is evidence that the immune system plays an important role in the pathogenesis of mental disorders and might be a source for diagnostic biomarkers relevant for the treatment of addictive disorders and associated psychiatric diseases (Fox et al., 2012; Araos et al., 2015; García-Marchena et al., 2017a; García-Marchena et al., 2017b). In the present study, we attempted to establish potential biomarkers of cocaine consumption and/or comorbid mental disorders by identifying certain, cytokines, chemokines and growth factors in the plasma of patients with these disorders. Previous studies (Araos et al., 2015; Pedraz et al., 2015) have revealed that the circulatory/immune system-related signal or growth factors are associated with CUD. The present results extend these observations and indicate that IL-17α, MIP-1α and TGFα are altered in subjects diagnosed with CUD during abstinence, although there was no association with the cocaine symptom severity, and that TGFα levels are also influenced by the presence of dual diagnosis. Although certain studies revealed positive associations between cytokines, chemokines, circulating growth factors and CUD (Sáez et al., 2011; Levandowski et al., 2016; Scherer et al., 2016), other studies reported a decreased expression of inflammatory proteins in CUD patients.

For instance, cocaine abusers were found to have decreased IL-10 levels compared with the levels found in social drinkers (Fox et al., 2012), and cocaine-dependent volunteers showed a decrease in the expression of cytokines such as TNFα and IL-6 (Irwin et al., 2007). Therefore, the role of these inflammatory signals in the neurobiology of addiction is not yet fully understood.

Cytokines and other inflammatory mediators are produced by several types of cells, mainly immune cells (e.g., lymphocytes and macrophages), and act through receptors in the regulation of crucial processes such as inflammation and embryogenesis. Some of these signals can cross the blood brain barrier and produce an inflammatory state that has been linked to dysfunctions on ascending monoaminergic systems (Petrulli et al., 2017). In fact, cytokines and their receptors are expressed in microglia, which are immune cells present in the brain that participate in processes such as remodeling and synaptic pruning (Paolicelli et al., 2011), as well as in astrocytes and neurons.

The decreased plasma levels of IL-17α, MIP-1α and TGFα are similar to that described in the same population for other cytokines and chemokines such as TNFα, MCP-1 and SDF-1 (Araos et al., 2015). Although cocaine has been described to produce a pro-inflammatory state, the decrease in the levels of IL-17α, MIP-1α and TGFα can be interpreted as a compensatory effect that reduces the cocaine-induced inflammatory tone along the period of abstinence in comparison with control subjects. A similar phenomenon has
been described for certain growth factor in our outpatient cohort of cocaine patients (Pedraz et al., 2015). However, none of the proteins that were analyzed in the present study were associated with CUD severity, unlike our previously described findings with IL-1β, fractalkine or SDF-1 as severity-sensitive factors (Araos et al., 2015).

As currently described in the literature, IL-17α plays a key role in autoimmune diseases in the CNS, such as multiple sclerosis (Zimmermann et al., 2017). However, there are no studies showing that IL-17α is involved in the effects of cocaine or other abused drugs. Regarding MIP-1α, this chemokine contributes to the modulation of immune responses, and MIP-1α is believed to be important in the pathogenesis of autoimmune and infectious diseases, as well as cancer (Snyder-Cappione et al., 2010). Recently, we have described that concentrations of MIP-1α are mildly affected in patients with alcohol use disorders (García-Marchena et al., 2017a). Although there is no scientific evidence linking this chemokine to the neuropharmacology of cocaine, it has been described that cocaine use activates the release of MIP-1α and promotes opening of the blood brain barrier, facilitating not only neuroinflammation but also viral infection from drug injectors (Zhang et al., 1998).

Considering that neurobiological changes associated with peripheral inflammatory states facilitate the action of psychostimulants (Petrulli et al., 2017), this clear association between MIP-1α and cocaine use demands further neurobiological studies.

It has been discovered that TGFα is related to a number of diseases. This ligand of the epidermal growth factor receptor has been proposed as a prognostic biomarker for gastric carcinoma (Fanelli et al., 2012), also for melanoma (Tarhini et al., 2014). Concerning CNS, TGFα is related to neurogenesis (Cooper & Isacson, 2004) and the decrease of TGFα in the plasma of cocaine patients might be related to a decrease in neurogenesis, which has been reported after acute administration of cocaine in animal models (Blanco-Calvo et al., 2014). This inhibition of neurogenesis might be associated with persistent rewarding memories for cocaine, and therefore, a decreased expression of TGFα might facilitate the persistence of cocaine abuse (Deschaux et al., 2014). Additionally, genetic models lacking TGFα in the brain result in super sensitivity to the psychostimulant effect of cocaine in a manner similar to that described for inflammation (Stanwood & Levitt, 2007). However, although a study in a population with heroin abuse disorders shows elevated levels of TGFα in injecting drug users with active consumption (Piepenbrink et al., 2016), to our knowledge, there are no studies relating TGFα to CUD.

Over the last decade, a growing number of studies have explored the potential role of cytokines, chemokines and growth factors in populations with mental disorders (Raison, Capuron & Miller, 2006; Ogłodek et al., 2015; Van Varsseveld et al., 2015; Engler et al., 2017; Michopoulos et al., 2017; Notter et al., 2017) and/or drug addiction, including CUD (Irwin et al., 2007; Parikh et al., 2014; Levandowski et al., 2016). Thus, certain studies have found reduced circulating cytokines in patients with generalized anxiety before and after Mindfulness treatment, when compared to their controls (Hoge et al., in press). Plasma levels of IL-17 and IL-23 are altered in schizophrenic patients relative to the levels in their controls (Li et al., 2016). Additionally, longitudinal studies described how personality traits predict IL-6 levels (Turiano et al., 2013).
An increase in IL-1β levels has been observed in patients with CUD with dual diagnosis compared to patients with CUD without a dual diagnosis (Araos et al., 2015). We believe that, together with monitoring cytokines, chemokines and growth factors, it is also necessary to characterize these patients with and without dual diagnosis, since they are patients with different therapeutic needs and approaches (Balhara, Kuppili & Gupta, 2017).

Related to the above, TGFα emerges as the only signal associated with both CUD and dual diagnosis. To our knowledge, there is no scientific literature on the relationship between CUD, dual diagnosis and plasma levels of TGFα.

In any case, the evaluation and validation of TGFα as a suitable candidate for a potential biomarker of both CUD and dual diagnosis could open new research lines for understanding the complexities of cocaine addiction and the associated dual diagnosis.

The evaluation of cytokines, chemokines and growth factors in plasma could improve the stratification of CUD patients undergoing treatment and complement therapeutic interventions, including the high risk of dual diagnosis. Additional studies would be needed to examine new molecules of the immune system in order to elucidate their role in the etiology of CUD.

Several limitations should be considered when discussing the present findings. First, larger studies are needed because variability is common when considering the association of immune signals with mental disorders. As an example, a meta-analysis of depressive patients with matched controls and their relationship with circulating levels of plasma cytokines shows that there is a considerable heterogeneity of results (Köhler et al., 2017). Second, it would also be necessary to include samples of depressed or anxious patients with no diagnosis of substance use disorders throughout life in order to compare CUD populations with or without dual diagnosis. Third, the small number of female patients is an important limitation of the present study. It would be necessary to increase the sample population of women in order to have a more comprehensive view of the results obtained. Fourth, it would be interesting to include in future studies subjects with active cocaine consumption to observe the effects of the presence of cocaine at circulating levels. It would also be necessary to control the pathway of confounding pharmacological treatment; as such, controlling the medical use of anti-depressants, anxiolytics and anti-psychotics would give more consistency to the current findings.

Finally, there is a need for integrating all the information concerning this multiplicity of inflammatory signals in a single model of cocaine addiction. Although certain factors such as TGFα might contribute to important aspects of cocaine addiction and associated psychiatric comorbidities, the complexity of the interactions of these signaling inflammatory proteins falls beyond our current understanding. Further basic and clinical research is needed to elucidate the role of TGFα and other factors in the pathogenesis of addiction and their utility as potential clinical biomarkers.

We conclude from this study that TGFα could be a potential biomarker of CUD and dual diagnosis in abstinent patients; moreover, additional studies are needed to investigate and validate the effects of TGFα in patients with CUD.
ADDITIONAL INFORMATION AND DECLARATIONS

Funding
This work was supported by RETICS Red de Trastornos Adictivos (RD12/0028/0021; RD16/0017/0001) funded by Instituto de Salud Carlos III (ISC-III) and European Regional Development Funds-European Union (ERDF-EU); Research projects funded by Ministerio de Economía y Competitividad and ISC-III (PI13/02261 and PI16/01953); Research projects funded by Ministerio de Sanidad, Servicios Sociales e Igualdad and Plan Nacional sobre Drogas (049/2009 and 049/2013); Research project funded by Consejería de Economía, Innovación y Ciencia, Junta de Andalucía and ERDF-EU (CTS-433); Research projects funded by Consejería de Salud y Bienestar Social, Junta Andalucía (PI0228-2013 and PI0823-2012). Antonia Serrano and Francisco Javier Pavón hold Miguel Servet research contracts funded by ISC-III and ERDF-EU (CP14/00173 and CP14/00212, respectively). Pablo Romero-Sanchiz holds a ‘Río Hortega’ research contract funded by ISC-III and ERDFEU (CM13/0115). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Grant Disclosures
The following grant information was disclosed by the authors:
Instituto de Salud Carlos III (ISC-III) and European Regional Development Funds-European Union (ERDF-EU): RD12/0028/0021, RD16/0017/0001.
Ministerio de Economía y Competitividad and ISC-III: PI13/02261, PI16/01953.
Ministerio de Sanidad, Servicios Sociales e Igualdad and Plan Nacional sobre Drogas: 049/2009, 049/2013.
Consejería de Economía, Innovación y Ciencia, Junta de Andalucía and ERDF-EU: CTS-433.
Consejería de Salud y Bienestar Social, Junta Andalucía: PI0228-2013, PI0823-2012.
ISC-III and ERDF-EU: CP14/00173, CP14/00212.
ISC-III and ERDFEU: CM13/0115.

Competing Interests
The authors declare there are no competing interests.

Author Contributions
• Rosa Maza-Quiroga analyzed the data, prepared figures and/or tables.
• Nuria García-Marchena performed the experiments, wrote the paper.
• Pablo Romero-Sanchiz, María Pedraz and Raquel Nogueira-Arjona performed the experiments.
• Vicente Barrios contributed reagents/materials/analysis tools.
• Antonia Serrano reviewed drafts of the paper.
• Juan Jesus Ruiz, Maribel Soria and Rafael Campos recruitment.
• Julie Ann Chowen and Jesus Argente contributed reagents/materials/analysis tools.
• Marta Torrens, Meritxell López-Gallardo and Eva María Marco reviewed drafts of the paper.
• Fernando Rodríguez de Fonseca conceived and designed the experiments.
• Francisco Javier Pavón and Pedro Araos conceived and designed the experiments, analyzed the data, wrote the paper, prepared figures and/or tables.

Human Ethics
The following information was supplied relating to ethical approvals (i.e., approving body and any reference numbers):

The study and protocols for recruitment were approved by the Ethics Committee of the CEI Provincial de Málaga in accordance with the ‘Ethical Principles for Medical Research Involving Human Subjects’ that was adopted in the Declaration of Helsinki by the World Medical Association (64th WMA General Assembly, Fortaleza, Brazil, October 2013). The data were protected by the Recommendation No. R (97) 5 of the Committee of Ministers to Member States on the Protection of Medical Data (1997) and the Spanish data protection act (Ley Orgánica 15/1999 de Protección de Datos, LOPD).

Data Availability
The following information was supplied regarding data availability:

The raw data has been provided as a Supplemental File.

Supplemental Information
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.3926#supplemental-information.

REFERENCES
Álvarez Y, Pérez-Mañá C, Torrens M, Farré M. 2013. Antipsychotic drugs in cocaine dependence: a systematic review and meta-analysis. Journal of Substance Abuse Treatment 45(1):1–10 DOI 10.1016/j.jsat.2012.12.013.

Araos P, Pedraz M, Serrano A, Lucena M, Barrios V, García-Marchena N, Campos-Cloute R, Ruiz JJ, Romero P, Suarez J, Baixeras E, De La Torre R, Montesinos J, Guerri C, Rodriguez-Arias M, Miñarro J, Martínez-Riera R, Torrens M, Chowen JA, Argente J, Mason BJ, Pavon FJ, Fonseca De Rodriguez, F. 2015. Plasma profile of proinflammatory cytokines and chemokines in cocaine users under outpatient treatment: influence of cocaine symptom severity and psychiatric co-morbidity. Addiction Biology 20(4):756–772 DOI 10.1111/adb.12156.

Balhara YP, Kuppili PP, Gupta R. 2017. Neurobiology of comorbid substance use disorders and psychiatric disorders: current state of evidence. Journal of Addictions Nursing 28(1):11–26 DOI 10.1097/JAN.0000000000000155.

Blanco-Calvo E, Rivera P, Arrabal S, Vargas A, Pavón FJ, Serrano A, Castilla-Ortega E, Galeano P, Rubio L, Suárez J, De Fonseca FR. 2014. Pharmacological blockade of either cannabinoid CB1 or CB2 receptors prevents both cocaine-induced conditioned locomotion and cocaine-induced reduction of cell proliferation in the hippocampus of adult male rat. Frontiers in Integrative Neuroscience 8(7):Article 106 DOI 10.3389/fnint.2013.00106.
Castilla-Ortega E, Serrano A, Blanco E, Araos P, Suárez J, Pavón FJ, Rodríguez de Fonseca F, Santín LJ. 2016. A place for the hippocampus in the cocaine addiction circuit: potential roles for adult hippocampal neurogenesis. Neuroscience & Biobehavioral Reviews 66:15–32 DOI 10.1016/j.neubiorev.2016.03.030.

Cearley CN, Blindheim K, Sorg BA, Krueger JM, Churchill L. 2011. Acute cocaine increases interleukin-1beta mRNA and immunoreactive cells in the cortex and nucleus accumbens. Neurochemical Research 36:686–692 DOI 10.1007/s11064-011-0410-9.

Chahua M, Sánchez-Niubó A, Torrens M, Sordo L, Bravo MJ, Brugal MT, Domingo-Salvany A. 2015. Quality of life in a community sample of young cocaine and/or heroin users: the role of mental disorders. Quality of Life Research 24(9):2129–2137 DOI 10.1007/s11136-015-0943-5.

Chibungu A, Gundareddy V, Wright SM, Nwabuo C, Bollampally P, Landis R, Eid SM. 2016. Management of cocaine-induced myocardial infarction: 4-year experience at an urban medical center. Southern Medical Journal 109(3):185–190 DOI 10.14423/SMJ.0000000000000430.

Coller JK, Hutchinson MR. 2012. Implications of central immune signaling caused by drugs of abuse: mechanisms, mediators and new therapeutic approaches for prediction and treatment of drug dependence. Pharmacology & Therapeutics 134(2):219–245 DOI 10.1016/j.pharmthera.2012.01.008.

Cooper O, Isacson O. 2004. Intrastriatal transforming growth factor α delivery to a model of Parkinson’s disease induces proliferation and migration of endogenous adult neural progenitor cells without differentiation into dopaminergic neurons. Journal of Neuroscience 24(41):8924–8931 DOI 10.1523/JNEUROSCI.2344-04.2004.

Cui C, Shurtleff D, Harris RA. 2014. Neuroimmune mechanisms of alcohol and drug addiction. International Review of Neurobiology 118:1–12 DOI 10.1016/B978-0-12-801284-0.00001-4.

Deschaux O, Vendruscolo LF, Schlosburg JE, Diaz-Aguilar L, Yuan CJ, Sobieraj JC, George O. Koob, GF, Mandyam CD. 2014. Hippocampal neurogenesis protects against cocaine-primed relapse. Addiction Biology 19(4):562–574 DOI 10.1111/adb.12019.

Engler H, Brendt P, Wischermann J, Wegner A, Röhling R, Schoemberg T, Meyer U, Gold R, Peters J, Benson S, Schedlowski M. 2017. Selective increase of cerebrospinal fluid IL-6 during experimental systemic inflammation in humans: association with depressive symptoms. Molecular Psychiatry 22:1448–1454 DOI 10.1038/mp.2016.264.

Fanelli MF, Chinen LTD, Begnami MD, Costa WL, Fregnami JHT, Soares FA, Montagnini AL. 2012. The influence of transforming growth factor-α, cyclooxygenase-2, matrix metalloproteinase (MMP)-7, MMP-9 and CXCR4 proteins involved in epithelial–mesenchymal transition on overall survival of patients with gastric cancer. Histopathology 61(2):153–161 DOI 10.1111/j.1365-2559.2011.04139.x.

Fox HC, D’Sa C, Kimmerling A, Siedlarz KM, Tuit KL, Stowe R, Sinha R. 2012. Immune system inflammation in cocaine dependent individuals: implications for medications development. Human Psychopharmacology 27(2):156–166 DOI 10.1002/hup.1251.
García-Marchena N, Araos P, Barrios V, Sánchez-Marin L, Chowen JA, Pedraz M, Castilla-Ortega E, Romero-Sanchiz P, Ponce G, Gavito AL, Decara J, Silva D, Torrens M, Argente J, Rubio G, Serrano A, Rodríguez de Fonseca F, Pavón FJ. 2017a. Plasma chemokines in patients with alcohol use disorders: association of CCL11 (Eotaxin-1) with psychiatric comorbidity. Front Psychiatry 7:Article 214 DOI 10.3389/fpsyt.2016.00214.

García-Marchena N, Araos P, Pavón FJ, Ponce G, Pedraz M, Serrano A, Arias F, Romero-Sanchiz P, Suárez J, Pastor A, De la Torre R, Torrens M, Rubio G, Rodríguez de Fonseca F. 2017b. Psychiatric comorbidity and plasma levels of 2-acyl-glycerols in outpatient treatment alcohol users. Analysis of gender differences. Adicciones 29(2):83–96 DOI 10.20882/adicciones.728.

Goldstein BI, Kemp DE, Soczynska JK, McIntyre RS. 2009. Inflammation and the phenomenology, pathophysiology, comorbidity, and treatment of bipolar disorder: a systematic review of the literature. The Journal of Clinical Psychiatry 70(8):1078–1090 DOI 10.4088/JCP.08r04505.

González-Saiz F, Vergara-Moragues E, Verdejo-García A, Fernández-Calderón F, Lozano OM. 2014. Impact of psychiatric comorbidity on the in-treatment outcomes of cocaine-dependent patients in therapeutic communities. Substance Abuse 35(2):133–140 DOI 10.1080/08897077.2013.812544.

Hasin DS, Trautman KD, Miele GM, Samet S. 1996. Psychiatric Research Interview for Substance and Mental Disorders (PRISM): reliability for substance abusers. The American Journal of Psychiatry 153(9):1195–1201 DOI 10.1176/ajp.153.9.1195.

Hoge EA, Bui E, Palitz SA, Schwarz NR, Owens ME, Johnston JM, Pollack MH, Simon NM. 2017. The effect of mindfulness meditation training on biological acute stress responses in generalized anxiety disorder. Psychiatry Research In Press DOI 10.1016/j.psychres.2017.01.006.

Irwin MR, Miller AH. 2007. Depressive disorders and immunity: 20 years of progress and discovery. Brain, Behavior, and Immunity 21(4):374–383 DOI 10.1016/j.bbi.2007.01.010.

Irwin MR, Olmos L, Wang M, Valladares EM, Motivala SJ, Fong T, Cole SW. 2007. Cocaine dependence and acute cocaine induce decreases of monocyte proinflammatory cytokine expression across the diurnal period: autonomic mechanisms. Journal of Pharmacology and Experimental Therapeutics 320(2):507–515 DOI 10.1124/jpet.106.112797.

Köhler CA, Freitas TH, Maes M, De Andrade NQ, Liu CS, Fernandes BS, Stubbs B, Solmi M, Veronese N, Herrmann N, Raison CL, Miller BJ, Lanctôt KL, Carvalho AF. 2017. Peripheral cytokine and chemokine alterations in depression: a meta-analysis of 82 studies. Acta Psychiatrica Scandinavica 135(5):373–387 DOI 10.1111/acps.12698.

Levandowski ML, Viola TW, Prado CH, Wieck A, Bauer ME, Brietzke E, Grassi-Oliveira R. 2016. Distinct behavioral and immunoendocrine parameters during crack cocaine abstinence in women reporting childhood abuse and neglect. Drug and Alcohol Dependence 167:140–148 DOI 10.1016/j.drugalcdep.2016.08.010.
Li H, Zhang Q, Li N, Wang F, Xiang H, Zhang Z, Su Y, Huang Y, Zhang S, Zhao G, Zhou R, Mao L, Lin Z, Cai W, Fang Y, Xie B, Zhao M, Hong W. 2016. Plasma levels of Th17-related cytokines and complement C3 correlated with aggressive behavior in patients with schizophrenia. Psychiatry Research 246:700–706 DOI 10.1016/j.psychres.2016.10.061.

Mestre-Pintó JI, Domingo-Salvany A, Martin-Santos R, Torrens M, PsyCoBarcelona Group. 2014. Dual diagnosis screening interview to identify psychiatric comorbidity in substance users: development and validation of a brief instrument. European Addiction Research 20(1):41–48 DOI 10.1159/000351519.

Michopoulos V, Powers A, Gillespie CF, Ressler KJ, Jovanovic T. 2017. Inflammation in fear and anxiety-based disorders: PTSD, GAD, and beyond. Neuropsychopharmacology 42(1):254–270 DOI 10.1038/npp.2016.146.

Moreira FP, Medeiros JRC, Lhullier AC, De Mattos Souza LD, Jansen K, Portela LV, Lara DR, Da Silva RA, Wiener CD, Oses JP. 2016. Cocaine abuse and effects in the serum levels of cytokines IL-6 and IL-10. Drug and Alcohol Dependence 158:181–185 DOI 10.1016/j.drugalcdep.2015.11.024.

Morgello S, Holzer 3rd CE, Ryan E, Young C, Naseer M, Castellon SA, Frol AB, Atkinson JH, Gelman BB, Grant I, Singer EJ. 2006. Interrater reliability of the Psychiatric Research Interview for Substance and Mental Disorders in an HIV-infected cohort: experience of the National NeuroAIDS Tissue Consortium. International Journal of Methods in Psychiatric Research 15(3):131–138 DOI 10.1002/mpr.189.

Neupane SP. 2016. Neuroimmune interface in the comorbidity between alcohol use disorder and major depression. Frontiers in Immunology 7:Article 625 DOI 10.3389/fimmu.2016.00655.

Notter T, Coughlin JM, Gschwind T, Weber-Stadlbauer U, Wang Y, Kassiu M, Vernon AC, Benke D, Pomper MG, Sawa A, Meyer U. 2017. Translational evaluation of translocator protein as a marker of neuroinflammation in schizophrenia. Molecular Psychiatry Epub ahead of print Jan 17 2017 DOI 10.1038/mp.2016.248.

Ogłodek EA, Szota AM, Just MJ, Moś DM, Araszkiewicz A. 2015. The MCP-1, CCL-5 and SDF-1 chemokines as pro-inflammatory markers in generalized anxiety disorder and personality disorders. Pharmacological Reports 67(1):85–89 DOI 10.1016/j.pharep.2014.08.006.

Paolicelli RC, Bolasco G, Pagani F, Maggi I, Sciami M, Panzanelli P, Giustetto M, Ferreira TA, Guiducci E, Dumas L, Ragozzino D, Gross CT. 2011. Synaptic pruning by microglia is necessary for normal brain development. Science 333(6048):1456–1458 DOI 10.1126/science.1202529.

Parikh N, Dampier W, Feng R, Passic SR, Zhong W, Frantz B, Blakey B, Aiamkitsumrit B, Pirrone V, Nonnemacher MR, Jacobson JM, Wigdahl B. 2014. Cocaine alters cytokine profiles in HIV-1-infected African American individuals in the DrexelMed HIV/AIDS genetic analysis cohort. Journal of Acquired Immune Deficiency Syndromes 66(3):256–264 DOI 10.1097/QAI.0000000000000163.

Pavón FJ, Araos P, Pastor A, Calado M, Pedraz M, Campos-Cloute R, Ruiz JJ, Serrano A, Blanco E, Rivera P, Suárez J, Romero-Cuevas M, Pujadas M, Vergara-Moragues...
E, Gornemann I, Torrens M, De la Torre R, Rodríguez de Fonseca F. 2013. Evaluation of plasma-free endocannabinoids and their congeners in abstinent cocaine addicts seeking outpatient treatment: impact of psychiatric co-morbidity. *Addiction Biology* 18(6):955–969 DOI 10.1111/adb.12107.

Pedraz M, Martín-Velasco AI, García-Marchena N, Araos P, Serrano A, Romero-Sanchiz P, Suárez J, Castilla-Ortega E, Barrios V, Campos-Cloute R, Ruiz JJ, Torrens M, Chowen JA, Argente J, De la Torre R, Santín LJ, Villanúa MÁ, Rodríguez de Fonseca F, Pavón FJ. 2015. Plasma concentrations of BDNF and IGF-1 in abstinent cocaine users with high prevalence of substance use disorders: relationship to psychiatric comorbidity. *PLOS ONE* 10(3):e0118610 DOI 10.1371/journal.pone.0118610.

Petrulli JR, Kalish B, Nabulsi NB, Huang Y, Hannestad J, Morris ED. 2017. Systemic inflammation enhances stimulant-induced striatal dopamine elevation. *Translational Psychiatry* 7(3):e1076 DOI 10.1038/tp.2017.18.

Piepenbrink MS, Samuel M, Zheng B, Carter B, Fucile C, Bunce C, Kiebala M, Khan AA, Thakar J, Maggirwar SB, Morse D, Rosenberg AF, Haughey NJ, Valenti W, Keefer MC, Kobie JJ. 2016. Humoral dysregulation associated with increased systemic inflammation among injection heroin users. *PLOS ONE* 11(7):e0158641 DOI 10.1371/journal.pone.0158641.

Raison CL, Capuron L, Miller AH. 2006. Cytokines sing the blues: inflammation and the pathogenesis of depression. *Trends Immunology* 27:24–31 DOI 10.1016/j.it.2005.11.006.

Sáez CG, Olivares P, Pallavicini J, Panes O, Moreno N, Massardo T, Mezzano D, Pereira J. 2011. Increased number of circulating endothelial cells and plasma markers of endothelial damage in chronic cocaine users. *Thrombosis Research* 128(4):e18–e23 DOI 10.1016/j.thromres.2011.04.019.

Scherer JN, Schuch S, Ornell F, Sordi AO, Bristot G, Pfaffenseller B, Kapczinski F, Kessler FH, Fumagalli F, Pechansky F, Von Diemen L. 2016. High levels of brain-derived neurotrophic factor are associated with treatment adherence among crack-cocaine users. *Neuroscience Letters* 630:169–175 DOI 10.1016/j.neulet.2016.07.050.

Slavich GM. 2016. Life stress and health: a review of conceptual issues and recent findings. *Teaching of Psychology* 43(4):346–355 DOI 10.1177/0739986316662768.

Snyder-Cappione JE, Tincati C, Eccles-James IG, Cappione AJ, Ndhllovu LC, Koth LL, Nixon DF. 2010. A comprehensive ex vivo functional analysis of human NKT cells reveals production of MIP1-α and MIP1-β, a lack of IL-17, and a Th1-bias in males. *PLOS ONE* 5(11):e15412 DOI 10.1371/journal.pone.0015412.

Stanwood GD, Levitt P. 2007. Waved-1 mutant mice are hypersensitive to the locomotor actions of cocaine. *Synapse* 61(4):259–262 DOI 10.1002/syn.20364.

Stertz L, Magalhaes PV, Kapczinski F. 2013. Is bipolar disorder an inflammatory condition? The relevance of microglial activation. *Current Opinion in Psychiatry* 26(1):19–26 DOI 10.1097/YCO.0b013e32835aad4b4.

Tarhini AA, Lin Y, Yeku O, La Framboise WA, Ashraf M, Sander C, Lee S, Kirkwood JM. 2014. A four-marker signature of TNF-RII, TGF-α, TIMP-1 and CRP is
prognostic of worse survival in high-risk surgically resected melanoma. *Journal of Translational Medicine* 12:Article 19 DOI 10.1186/1479-5876-12-19.

**Tomasi D, Wang GJ, Wang R, Caparelli EC, Logan J, Volkow ND. 2015.** Overlapping patterns of brain activation to food and cocaine cues in cocaine abusers. *Human Brain Mapping* 36(1):120–136 DOI 10.1002/hbm.22617.

**Torrens M, Gilchrist G, Domingo-Salvany A, PsyCoBarcelona Group. 2011.** Psychiatric comorbidity in illicit drug users: substance-induced versus independent disorders. *Drug and Alcohol Dependence* 113:147–156 DOI 10.1016/j.drugalcdep.2010.07.013.

**Torrens M, Serrano D, Astals M, Pérez-Domínguez G, Martín-Santos R. 2004.** Diagnosing comorbid psychiatric disorders in substance abusers: validity of the Spanish versions of the Psychiatric Research Interview for Substance and Mental Disorders and the Structured Clinical Interview for DSM-IV. *American Journal of Psychiatry* 161(7):1231–1237 DOI 10.3109/10826084.2012.663296.

**Turiano NA, Mroczek DK, Moynihan J, Chapman BP. 2013.** Big 5 personality traits and interleukin-6: evidence for “healthy Neuroticism” in a US population sample. *Brain, Behavior, and Immunity* 28:83–89 DOI 10.1016/j.bbi.2012.10.020.

**Van Varsseveld NC, Van Bunderen CC, Sohl E, Comijs HC, Penninx BW, Lips P, Drent ML. 2015.** Serum insulin-like growth factor 1 and late-life depression: a population-based study. *Psychoneuroendocrinology* 54:31–40 DOI 10.1016/j.psyneuen.2015.01.014.

**Verdejo-Garcia A, Verdejo-Román J, Albein-Urios N, Martinez-González JM, Soriano-Mas C. 2015.** Brain substrates of social decision-making in dual diagnosis: cocaine dependence and personality disorders. *Addiction Biology* 22(2):457–467 DOI 10.1111/adb.12318.

**Vergara-Moragues E, González-Saiz F, Lozano OM, Betanzos Espinosa P, Fernández Calderón F, Bilbao-Acebos I, Pérez García M, Verdejo Garcia A. 2012.** Psychiatric comorbidity in cocaine users treated in therapeutic community: substance-induced versus independent disorders. *Psychiatry Research* 200(2–3):734–741 DOI 10.1016/j.psychres.2012.07.043.

**Zhang L, Looney D, Taub D, Chang SL, Way D, Witte MH, Graves MC, Fiala M. 1998.** Cocaine opens the blood-brain barrier to HIV-1 invasion. *Journal of Neurovirology* 4(6):619–626 DOI 10.3109/13550289809114228.

**Zimmermann J, Emrich M, Krauthausen M, Saxe S, Nitsch L, Heneka MT, Campbell IL, Müller M. 2017.** IL-17A promotes granulocyte infiltration, myelin loss, microglia activation, and behavioral deficits during cuprizone-induced demyelination. *Molecular Neurobiology* Epub ahead of print Jan 13 2017 DOI 10.1007/s12035-016-0368-3.