FAAH genotype, CRFR1 genotype, and cortisol interact to predict anxiety in an aging, rural Hispanic population: A Project FRONTIER study

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**ABSTRACT**

The neurophysiological underpinnings involved in susceptibility to and maintenance of anxiety are not entirely known. However, two stress-responsive systems, the hypothalamic-pituitary-adrenal axis and the endocannabinoid system, may interact in anxiety. Here, we examine the relationship between FAAH genotype, CRFR1 genotype, baseline cortisol, and state anxiety in a rural adult population using data from Project FRONTIER. We predicted that FAAH A (AA and AC vs. CC; rs324420) and three CRFR1 SNP minor alleles (rs7209436 C $\rightarrow$ T [minor allele]; rs110402, G $\rightarrow$ A [minor]; and rs242924 G $\rightarrow$ T [minor]), would interact to predict low baseline cortisol and low state anxiety scores. We found partial support for our prediction. In CRFR1 minor carriers, the FAAH AA or AC (vs. CC) genotype was associated with higher cortisol and with lower anxiety. In CRFR1 non-minors, those with FAAH AA or AC (vs. CC) showed decreased cortisol and higher anxiety. These results suggest that FAAH CC genotype only conveys risk for anxiety in individuals who are also carriers of the CRFR1 minor combination. FAAH genotype was significantly associated with baseline cortisol but was not independently associated with anxiety. Contrary to our predictions, baseline cortisol was negatively associated with anxiety. Lastly, we did not find any independent relationships between any of our SNPs and baseline cortisol or anxiety. These data suggest FAAH and cortisol interact to predict state anxiety, but that the relationship depends on CRFR1 genotype. The Project FRONTIER dataset is supported by Texas Tech University Health Sciences Center Garrison Institute on Aging.

1. Introduction

According to National Institute of Mental Health (NIMH), over 40 million adults in the US, or roughly 18% of the population, suffer from anxiety, and nearly 30% of adults will experience an anxiety disorder during their lifetime (National Institute of Mental Health, 2017; Anxiety and Depression Association of America). Despite its prevalence, the neurobiological underpinnings of anxiety onset and maintenance are not fully understood. Using the NIMH Research Domain Criteria (RDoC) approach (Cuthbert and Insel, 2013), recent studies using humans and animal models have determined various biomarkers specific to anxiety. The RDoC aims to incorporate many dimensions including behavior, genetics, neurobiology, environment, and experiential effects into diagnosis and categorization of mental disorders, with the goal that doing so will aid in understanding disease etiology and will enhance treatment options (Insel, 2014) also see [Holmes and Patrick, 2018]). Due to the reliance on fear and threat in the diagnosis of anxiety disorders, studies have focused on the physiological systems associated with response to threats (stressors), specifically the hypothalamic-pituitary-adrenal (HPA) axis and the endocannabinoid system (Clinchy et al., 2011; Harris and Carr, 2016; Lang et al., 2016; McEwen et al., 2015; Perusini and Fanselow, 2015). Data suggest these two systems play both independent and interacting roles in the onset and maintenance of anxiety.

Dysregulation of the HPA axis, resulting in alteration to both baseline and stress-induced glucocorticoid levels (e.g., cortisol and corticosterone), has been implicated in several diseases and psychopathologies, including anxiety (McEwen et al., 2015, 2016; Shin and...
with a threonine (C385A, rs324420), alters the regulation of FAAH, but pact anxiety. Specifically, a substitution of a proline at amino acid 129 with a threonine (C385A, rs324420), alters the regulation of FAAH, but not its enzymatic activity, making it more vulnerable to proteolytic degradation (Chiang et al., 2004; Sipe et al., 2002). The AA or AC genotype is associated with decreased FAAH, increased AEA levels, and reduced amygdala activity (increased inhibitory tone) comparison to the CC genotype (Gunduz-Cinar et al., 2013). Thus, CC individuals seem to be at higher risk for anxiety disorders and anxious behavior as compared to A carriers, and this effect has been shown in humans and in a knock-in mouse model (Dincheva et al., 2015; Hariri et al., 2009). Given these findings, FAAH has recently been tagged as a possible therapeutic target for anxiety disorders (Hill et al., 2013a; Patel et al., 2017).

When taken together, data suggest CRF plays a central role in linking the HPA axis, the eCB system, and the amygdala with anxiety (Hill et al., 2009, 2010a; Hillard et al., 2011). In elaborate studies in mice, Gray and colleagues (Gray et al., 2015) showed stress-associated increase in CRF concentration in the amygdala resulted in CRFR1 activation, an increase in FAAH activity, and thus decrease AEA levels. Overall, this cascade results in decreased AEA binding to CB1 receptors resulting in increased glutamate release, and increased activity of the basolateral amygdala resulting in increased expression of anxiety-like behaviors (Gray et al., 2015); this effect appeared to be independent of initial stress-related glucocorticoid increase. These data suggest one mechanism by which stress increases anxiety is via CRFR1 impacts on FAAH activity and AEA binding in the basolateral amygdala. Another recent study using a FAAH inhibitor in rats corroborates these findings (Natividad et al., 2017).

Given the relationships among the HPA axis, CRFR1, FAAH, and anxiety, we hypothesized that baseline cortisol and SNPs in CRFR1 and FAAH genes interact to influence state anxiety in humans (see Fig. 1). Here, we determined if baseline cortisol, FAAH, and CRFR1 SNPs are related to state anxiety levels, measured via the Beck Anxiety Inventory (BAI), in an aged, majority Hispanic, rural, community dwelling human population. We proposed a moderated mediation in that baseline cortisol mediates the relationship between FAAH SNP (rs324420) and state anxiety, and that effect is moderated by CRFR1 genotype. Specifically, we predict that FAAH A (AA and AC vs CC) and CRFR1 SNP minor alleles (AA, TT, TT) will interact to predict low baseline cortisol and those individuals will have the lowest state anxiety scores. Additionally, for comparison with published data on other sample populations, we determined if baseline cortisol and any of the four SNPs are individually associated with anxiety scores, and if any of the four SNPs are related to baseline cortisol. To date, only one study, to our knowledge, has investigated the interactive effects of CRFR1 (rs110402; AA vs. GG and GA) and FAAH (rs324420; CC vs. AC and AA) genotype on anxiety-related markers in humans (Demers et al., 2016). Contrary to our predictions based on previous data (see above), using fMRI data and a moderated mediation analysis, Demers and colleagues found that individuals high in AEA signaling (FAAH AA or AC) with CRFR1 protective allele (AA vs. AG and GA) had blunted basolateral amygdala habituation and increased risk of anxiety disorders (via the DSM-IV criteria); they did not assess cortisol. Thus, it is possible that we could find results predicted by the literature outlined above, or, we may corroborate findings of Demers and colleagues and find the opposite relationship.

2. Methods

2.1. Participants

Data were obtained from the Texas Tech University Health Sciences Project FRONTIER (Facing Rural Obstacles to healthcare Now through Intervention, Education, and Research) database (https://www.ttuhsc.edu/ruralhealth/researchgroup/frontier.aspx). Project FRONTIER is funded by Texas Tech University Health Sciences Center Garrison Institute on Aging and is currently ongoing. Study coordinators set-up appointments at study participant’s rural county hospital every 3 years.
for testing and data collection. All data are collected by Project FRONTIER personnel and archived on a secure computer or in biobank freezers (biological specimens). Whole blood is collected by a trained phlebotomist at each visit. All data are collected with patient consent and TTUHSC Institutional Review Board approval was obtained by Project FRONTIER coordinators. Authors of this study did not directly interact with any of the participants. Authors obtained the following de-identified samples or variable data for 193 individuals: frozen whole blood, frozen serum, Beck Anxiety Inventory scores, last grade level completed in school, sex, age, race (white/Caucasian, black, or other), and ethnicity (Hispanic or not Hispanic). Participants from this study were community dwelling at the time of data collection and as long as they were over 40, were not excluded from the study for any reason. The average age of entry into the study was 58.27 yrs (range 40–87). Data obtained for this study were from the second appointment (year 3 of study).

Demographic information for the 193 participants is as follows:

- 92.2% Caucasian, 50.5% Hispanic, and 72.5% female. Self-reported highest grade level completed averaged 10.1 (range: 0–19). The average age at interview and data collection for samples obtained for our analysis was 61.09 yrs (std dev. 11.66, range: 42–90). All procedures performed by the authors were approved by the Texas Tech University IRB.

2.2. Project FRONTIER Beck Anxiety Inventory (BAI)

The BAI is a commonly used 21-item, self-report questionnaire which asks participants to rank anxiety symptoms on their severity over the past week using a 0–3 point Likert scale (Beck et al., 1988; Beck and Steer, 1993). The test has high internal reliability and discriminant validity (Fydrich et al., 1992; Kabacoff et al., 1997). Scores for each item are added and possible total score ranges from 0 to 63 (max score of 3 on each of 21 items). Recommended clinical cut-offs are: 0–7 minimal anxiety, 8–15 mild anxiety, 16–25 moderate anxiety, over 26,
severe anxiety (Carney et al., 2011); clinically significant anxiety according to Beck and Steer (1993) is 16. For all participants, BAI scores averaged 4.94 (range: 0–34; n = 193) and were thus in the minimally anxious range. Only 11 participants scored over 16 with 5 of those being higher than 26. Thus, only a handful of participants involved were likely clinically anxious, allowing us to correlate genotypes with measures of state anxiety in a majority non-clinical, aged adult, rural population.

2.3. DNA extraction and SNP genotyping

Whole blood was used for DNA extraction. Extraction was done using a commercially available kit as per the manufacturer's instructions (FlexiGene kit # 51206, Qiagen), except for the following modification: extracted DNA was suspended in molecular grade water instead of the storage buffer provided with the Qiagen kit. Following extraction, concentration of DNA in each sample was determined by Nanodrop spectrophotometry (average: 222 ng/ul) and the 260/280 ratio was inspected for purity. Extracted DNA was diluted with molecular grade water to a concentration of 10 ng/ul, aliquoted, and frozen for future use.

Genotyping analysis was performed on an Applied Biosystems 7300 quantitative PCR machine using commercially available TaqMan allelic discrimination SNP genotyping assays (Applied Biosystems, ThermoFisher Scientific). Participant genotype for FAAH (rs324420 [A or C]; kit C1897306), and three CRFR1 loci (rs110402 [A or G], kit C2544843; rs7209436 [C or T], kit C1570087; rs242924 [T or G], kit C2257689) as per the kit manufacturer's instructions. Genotype data for 176–186 individuals were available for analysis (FAAH: 186; CRFR1: 176 [rs110402], 185 [rs7209436], and 181 [rs242924]).

2.4. Serum cortisol assay

Project FRONTIER personnel obtained serum samples from blood collected at each visit. Of the 193 participant datasets requested, 191 had serum samples available. Serum was collected from blood and frozen in aliquots in the biobank. All blood samples were collected between 7:30 a.m. and 2:45 p.m. (only two samples were collected after 12 noon), and 90% of samples were collected between 7:00 and 9:30 a.m., thus we used time of day as a covariate in analyses. Concentration of serum cortisol was determined using a commercially available radioimmunoassay kit (ImmucheM, 07–221102; MP Biomedicals, Solon, OH) according to manufacturer's instructions.

2.5. Statistical analysis

All analyses were completed using SPSS v. 22 and alpha was set at 0.05. Moderated mediation analyses (Preacher & Hayes, 2008; Preacher et al., 2007) were conducted to examine the conditional indirect effect of baseline cortisol on the interaction of FAAH_A (AA and AC vs CC) and CRF1_minor (minor at all 3 SNPs: AA, TT, TT) on anxiety (for the benefits of this approach see [Antonakis et al., 2010; Hayes, 2009]), controlling for time of day the blood was drawn. The sexes were analyzed together. For this analysis we used model 7 in the process macro for SPSS (Hayes, 2018) with 50,000 bootstrapped samples (for the benefits of bootstrapping see [Hayes and Preacher, 2010; MacKinnon et al., 2000]). Note, for the CRFR1 minor grouping, the total n is 185 as we were able to include all participants with known CRFR1 genotype because even if they were missing data for one SNP, the other two did not meet the CRFR1 minor criteria.

Additional analyses for association between SNP and anxiety or cortisol concentration were completed via one-way ANOVA; effect sizes are reported as partial eta squared. Anxiety and cortisol association was assessed using Pearson's r.

3. Results

3.1. Moderated mediation

The total effect of the interaction between FAAH_A and CRF1_minor on anxiety was not statistically significant (b = −0.890, SE = 0.941, 95% CI bias corrected [-2.75, 0.967]). Results from the moderated mediation analysis indicated that the total indirect effect (interaction between FAAH_A and CRF1_minor to cortisol to anxiety) was significant − b = −0.721, SE = 0.505, 95% CI bias corrected [-2.00, −0.016]. Examining this moderated mediation, the conditional indirect effect of cortisol on anxiety was positive for CRF1_minor carriers, b = 0.542, SE = 0.380, 95% CI bias corrected [-0.012, 1.43], meaning that in CRF1_minor carriers, having the FAAH CC (compared to AA, AC) genotype is associated with lower cortisol and here lower cortisol is associated with increased anxiety. The conditional indirect effect was negative for CRF1_non-minor carriers, b = −0.180, SE = 0.254, 95% CI bias corrected [−0.820, 0.248], meaning that in CRF1_non-minor carriers, having the FAAH CC (compared to AA, AC) genotype is associated with increased cortisol and here increased cortisol is associated with lowered anxiety. Overall, the moderated mediation results suggest that the moderation of cortisol between FAAH_A and anxiety goes in one direction for CRF1_minor carriers and goes in the opposite direction for CRF1_non-minors. A summary of predictions and results are presented in Table 1 and results from the model are presented in Fig. 2.

3.2. Genotypes and associations between SNPs, cortisol, and anxiety

Genotype data are shown in Table 2. Individual genotype did not impact baseline cortisol (Table 3) or scores on the Beck Anxiety Inventory (Table 4). Baseline cortisol and state anxiety were negatively correlated in our total sample: as cortisol decreased, anxiety increased (r = −0.175, p = 0.015, n = 191).

4. Discussion

We found partial support for our prediction that FAAH A (AA and AC vs CC) and CRF1 SNP minor alleles (AA, TT, TT), would interact to predict low baseline cortisol and low state anxiety scores. In CRF1 minor carriers, the FAAH AA or AC (vs. CC) genotype was associated with higher cortisol and with lower anxiety. Additionally, in CRF1 non-minors, those with FAAH AA or AC (vs. CC) showed decreased cortisol and higher anxiety. Based on data from this population, it seems the minor alleles of CRF1 and the AA or AC genotype of FAAH are related to reduced anxiety, but, surprisingly, related to higher, and not lower, baseline cortisol. Additionally, among individuals with the protective CRF1 minor combination, those with the FAAH CC genotype that confers risk had higher anxiety, but, those with the FAAH CC genotype and without the protective CRF1 minor combination had lower anxiety. Thus, it seems the FAAH CC genotype only confers risk for anxiety in individuals who are also carriers of the CRF1 minor

Table 1

| CRF1 Minor SNP carriers (“protective”, decreased HPA activity) | CRF1 non-minor SNP carriers (“more risky”, increased HPA activity) |
|------------------|------------------|
| FAAH AA/AC (High AEA) | FAAH AA/AC (High AEA) |
| FAAH CC (Low AEA) | FAAH CC (Low AEA) |
| Higher cortisol | Higher cortisol |
| Lower anxiety | Higher anxiety |
| Higher cortisol | Higher anxiety |

Table 2

| FAAH AA/AC | CRF1 Minor SNP carriers “protective”, decreased HPA activity | CRF1 non-minor SNP carriers “more risky”, increased HPA activity |
|------------------|------------------|------------------|
| FAAH CC (Low AEA) | FAAH AA/AC (High AEA) | FAAH AA/AC (High AEA) |
| Higher cortisol | Higher cortisol | Higher cortisol |
| Lower anxiety | Higher anxiety | Higher anxiety |

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When dissecting our model overall, FAAH genotype was significantly associated with baseline cortisol, in that CC individuals had higher cortisol than did AA and AC individuals, which is in line with our initial predictions. But, FAAH genotype was not independently associated with anxiety, and baseline cortisol was negatively associated with anxiety score; as cortisol decreased, anxiety increased, which is contrary to our prediction. Lastly, contrary to other studies we did not find any independent relationships between any SNPs and baseline cortisol or anxiety score.

Although unexpected based on our predictions, low baseline and post-stress cortisol being associated with high anxiety has been previously documented in both clinical and healthy populations of humans (Beaton et al., 2006; Jezova et al., 2004; Shirotsuki et al., 2009; Vingerhoets et al., 1996), and in rodents (Cohen et al., 2006). But, none of the above studies assess genotype. The reason for the association between decreased baseline and post-acute-stress cortisol and anxiety is not clear. However, recent studies exploring the interaction between the HPA axis and the endocannabinoid system may provide insight (Balsevich et al., 2017). For example, it is well documented (in rodents) that acute stress (likely via increased CRF) decreases the amount of AEA in the BLA resulting in increased anxiety and in HPA axis activation (Gray et al., 2015), but glucocorticoids can have different effects from stress. Exposure to elevated glucocorticoids without stress results in increased AEA in the amygdala of rats within 10 min (Hill et al., 2010b). Additionally, the site of sampling, for example brain vs. periphery, matters. Whereas in rodents, acute stress decreases amygdala AEA, in humans, acute psychosocial stress increased circulating AEA (Dlugos et al., 2012), and circulating cortisol and AEA have been shown to be positively correlated (Hill et al., 2013b). This AEA-enhancing effect of stress and glucocorticoids is thought to help reign in the initial response to stress (Morena et al., 2016), suggesting that individuals with lower baseline and post-acute-stress cortisol may have decreased AEA and thus more activity in the BLA. Moreover, circulating AEA is negatively correlated with anxiety in healthy and psychiatric populations – lower AEA is related to higher anxiety (Dlugos et al., 2012; Hill et al., 2008), suggesting that when taking this evidence into account lower cortisol could be associated with higher anxiety. But, individuals with lower baseline AEA tended to have higher post-stress cortisol, and baseline cortisol is not always correlated with baseline AEA (Dlugos et al., 2012), suggesting more information is needed. Glucocorticoids are necessary for sculpting the functional connectivity between the amygdala and prefrontal cortex and for normal fear extinction (Hill et al., 2018), and reduced cortisol, along with elevated CRF, is found in individuals with PTSD, a disorder characterized, at least in part, by

![Fig. 2. Cortisol mediated the interaction between FAAH genotype and CRFR1 minor on anxiety. Pathway coefficients represent unstandardized b values. The 95% confidence interval (95% CI) is presented below the b value, if the interval does not include zero then it is significant at p < 0.05. The dark dotted line is the mediation of cortisol on FAAH and anxiety for CRFR1 minor allele carriers and the gray dotted line is the mediation of cortisol on FAAH and anxiety for CRFR1 non-minor allele carriers.](image)

### Table 2
Genotype frequencies.

| Gene          | Restriction site | Genotype (n) | Total n |
|---------------|-----------------|--------------|---------|
| FAAH          | rs324420        | AA (20)      | 186     |
|               |                 | AC (74)      |         |
|               |                 | CC (92)      |         |
| CRFR1         | rs110402        | AA (64)      | 176     |
|               |                 | AG (74)      |         |
|               |                 | GG (38)      |         |
| CRFR1         | rs7209436       | TT (63)      | 185     |
|               |                 | CT (80)      |         |
|               |                 | CC (42)      |         |
| CRFR1 minor   | rs110402; rs7209436; rs242924 | AA/TT/TT (60) | 185 |
| CRFR1         | rs242924        | TT (60)      | 125     |

### Table 3
Relationship between genotype and baseline cortisol (ug/dl) for all participants (sexes combined). Cortisol data presented are means and standard error, statistical results are from ANOVA, effects size is partial eta squared.

| Gene          | Restriction site | Cortisol (genotype) | Main effect of genotype | Effect size |
|---------------|-----------------|---------------------|-------------------------|-------------|
| FAAH          | rs324420        | 18.90 ± 0.82 (AA/AC) | 18.38 ± 0.83 (CC)       | F 1,179 = 0.19, P = 0.660 0.001 |
| CRFR1         | rs110402        | 19.28 ± 1.00 (AA)   | 18.77 ± 0.94 (AG)       | F 2,171 = 1.38, P = 0.256 0.016 |
| CRFR1         | rs7209436       | 19.40 ± 1.00 (TT)   | 18.65 ± 0.89 (CT)       | F 2,180 = 1.23, P = 0.484 0.008 |
| CRFR1         | rs242924        | 19.24 ± 0.98 (TT)   | 19.45 ± 0.91 (TG)       | F 2,176 = 1.31, P = 0.272 0.015 |
| CRFR1 minor   | rs110402; rs7209436; rs242924 | 19.78 ± 0.72 (AA/TT/TT) | 18.02 ± 0.67 (AG/GG) | F 1,182 = 0.72, P = 0.015 0.006 |
| CRFR1         | rs110402        | 19.28 ± 0.76 (TT/GG) | 18.4 ± 0.76 (CT/CC)    | F 1,181 = 0.88, P = 0.349 0.005 |
| CRFR1         | rs242924        | 19.40 ± 0.97 (TT)   | 18.61 ± 0.73 (TG/GG)   | F 1,177 = 0.26, P = 0.608 0.002 |
overactivity of the amygdala and impaired fear processing (Hill et al., 2018). Taken together, these data suggest low cortisol can be associated with endocannabinoid signaling and anxiety.

It is important to note, however, that the bidirectional interactions between the endocannabinoid system and the HPA axis are complex (Balsevich et al., 2017; Morena et al., 2016) and involve not just the BLA and PVN, but several other brain regions (e.g., hippocampus, prefrontal cortex, ventral tegmental area, bed nucleus of the stria terminalis) and mediators (e.g., 2-AG, arginine vasopressin, GABA, 5-HT). Additionally, the time course of hormone elevation, time point of sample collection, location of sample collection, level of hormone (baseline vs. post-stress), duration of stress (acute vs. chronic), past life experiences (e.g., childhood or adult trauma), and species (e.g. rodents vs humans) can all impact outcomes (Balsevich et al., 2017; Hill et al., 2018; Morena et al., 2016). Less is known about the baseline interactions of these systems as more work has investigated impacts of acute or chronic stress and/or articular glucocorticoid elevation, and, due to ability to control variables and manipulate and analyze brains, rodents have been the primary study species. Future studies to illuminate more precisely how durations (acute vs. chronic) and species-specific levels (e.g., circadian peak or post-stress) of glucocorticoid increase alter amygdalar AEA and FAAH, and anxiety-like behavior are needed to determine the interacting role these systems play in anxiety. For our study, it is important to point out that no specific stressor was applied prior to collection of blood samples used for assay of cortisol, but it is possible that hormone levels were not true representations of baseline as some participants may have found anticipation of a medical exam as some participants may have found anticipation of a medical exam stressful. We did use time of day of blood sample as a covariate to control for circadian variation in cortisol. However, even if our samples are not true baseline values, our data are still supported by other studies showing baseline and post-stress cortisol are lower in those with increased anxiety. Additionally, we were not able to measure AEA in our study, but future studies should measure levels of circulating glucocorticoids and endocannabinoids in the same participants.

We found the FAAH CC genotype only confers risk for anxiety if the CRFR1 minor allele combination is also present. Additionally, the FAAH AA/AC genotype was riskier than the CC if the three CRFR1 minor alleles were not present. This result is somewhat contrary to our prediction as we hypothesized the FAAH AA/AC and the CRFR1 minor combination would buffer individuals from anxiety both in combination and on their own. While it is unknown why we found this flipped effect of FAAH SNP depending on CRFR1 SNP status, it may be that in response to CRF signaling (which may vary by genotype and/or stress), individuals with higher baseline AEA (those with AA/AC) experience a greater relative drop in AEA compared to the CC individuals that started with less AEA. If the greater drop corresponds to proportionally greater release of inhibition (by decreased binding at more CB1 receptors) it may therefore relate to more glutamatergic output and thus greater release of inhibition (by decreased binding at more CB1 receptors) in the BLA among individuals who differ in FAAH genotype is not available. However, data from rodents suggest that the level of CB1 binding is important for signaling - the higher the CB1 agonist concentration, the lower the firing rate of BLA neurons (Pistis et al., 2004), suggesting that a more pronounced decrease in AEA could possible result in greater BLA output. Additionally, recent work highlights the importance of CB1 in understanding the interactions among the EC system, stress, and anxiety. For example, chronic stress, sustained glucocorticoids, and early life stress can decrease CB1 receptor density in the brain; blocking CB1 can increase anxiety in baseline and post-stress scenarios and can increase HPA axis activity (Morena et al., 2016; Balsevich et al., 2017; Ibarra-Lecue et al., 2018; Hill et al., 2019). Moreover, another recent study found FAAH A carriers had higher anxiety (and depression) than CC individuals, but only if they were exposed to childhood adversity (Lazary et al., 2016); CRFR1 genotype, CB1, and cortisol were not assessed. These data suggest that the role of CB1 is likely important and may help to explain how gene by environment interactions impact anxiety. Future studies should determine if CRFR1 minor alleles play a role in the relationship between FAAH genotype, CB1 dynamics, and anxiety.

It is important to note that despite the abundance of data on the role of CRFR1 SNPs in various stress and affective state outcomes, it is unclear what the functional significance of these SNPs are at the molecular/cellular level (Binder and Nemeroff, 2010). The gene encoding CRFR1 is located on chromosome 17 and contains 13 exons spanning 51 kb (Binder and Nemeroff, 2010). The three CRFR1 SNPs included here are located near the 5′ end and are in the first and second intron region (Chen et al., 1993; Vita et al., 1993). But it is unclear what these SNPs do to CRFR1 functionality; future studies addressing this would be beneficial for interpreting data. Additionally, several previous reports of CRFR1 minor alleles being protective are in situations that involved early life trauma (Tyrka et al., 2009; Bradley et al., 2008; Polanczyk et al., 2009). Specifically, in white, non-Hispanic individuals (n = 129), those with the common homozygous genotypes of GG (vs. AG or AA; rs110402) or GG (vs. GT or TT; rs242924) who experienced childhood trauma showed increased cortisol response to the Dex/CRF challenge test; but, there was no impact of genotype on cortisol in individuals without childhood trauma (Tyrka et al., 2009). Moreover, the presence of two copies of the minor allele for each of the three SNPs was found to be protective against adult depression in African American, non-Hispanic individuals (n = 422) who experienced severe early life stress/truma (Bradley et al., 2008). Additionally, in an additive model, individuals (n = 368, Caucasians) homozygous for the minor allele at each SNP (TT, AA, or TT) showed decreased cortisol response to an acute stressor (Trier Social Stress Test; Mahon et al., 2013). And, higher baseline cortisol was associated with increased trait anxiety (Spielberger STAI-T) in individuals homozygous for the common allele of rs7209436 (CC) and rs110402 (GG); authors did not report main effects of genotype on anxiety measures (Mahon et al., 2013). Thus, it is therefore possible that a gene by environment interaction is important for interpreting the role of CRFR1 and FAAH genotype (see Lazary et al., 2016) in stress physiology and affect (Heim and Binder, 2012). In line with this, recent studies have shown that early life trauma can lead to differential DNA methylation (Roberts et al., 2018) and that this

Table 4

| Gene      | Restriction site | BAI (genotype) | Main effect of genotype | Effect size |
|-----------|-----------------|----------------|-------------------------|------------|
| FAAH      | rs324420        | 5.15 ± 1.45 (AA) | 5.55 ± .75 (AC)         | 4.47 ± 0.67 (CC) | F 2,183 = 0.59, P = 0.558 | 0.006 |
| CRFR1     | rs110402        | 4.72 ± 0.81 (AA) | 4.78 ± .75 (AG)         | 5.18 ± 1.05 (GG) | F 2,173 = 0.07, P = 0.935 | 0.001 |
| CRFR1     | rs242924        | 4.57 ± .80 (TT)  | 5.02 ± .72 (CT)         | 5.40 ± 0.99 (CC) | F 2,182 = 0.37, P = 0.694 | 0.004 |
| CRFR1 minor | rs110402 r7209436 r242924 | 4.40 ± 0.823 (AA/TT/TT) | 5.00 ± .75 (TG) | 5.24 ± 1.01 (GG) | F 2,176 = 0.15, P = 0.859 | 0.002 |
| CRFR1     | rs110402        | 4.72 ± 0.81 (AA) | 4.92 ± 0.61 (AG/GG)     | 5.18 ± 0.99 (CC) | F 2,184 ± 0.46, P = 0.498 | 0.003 |
| CRFR1     | rs7209436       | 4.37 ± 0.80 (TT) | 5.16 ± 0.58 (CT/CC)     | 5.18 ± 0.64 (CC) | F 1,183 = 0.64, P = 0.426 | 0.003 |
| CRFR1     | rs242924        | 4.57 ± 0.80 (TT) | 5.09 ± 0.60 (TG/GG)     | 5.18 ± 0.67 (CC) | F 1,179 = 0.27, P = 0.605 | 0.001 |
impact is directly related to stress-responsive genes (FKBP5) and affect, mainly depression (Binder, 2009; Binder et al., 2008; A. Zannis and Binder, 2014; Zannis et al., 2016). We do not have information on exposure to early life trauma in our study population, but this would be an interesting area for future study.

Overall, we partially confirmed our predictions, and, when compared to the other existing dataset exploring the interaction among CRFR1, FAAH, and anxiety, our data are somewhat confirmatory. Demers and colleagues (Demers et al., 2016) found one (protective) CRFR1 minor allele (rs110402) and the (less risky) FAAH AA or AC alleles were associated with increased anxiety disorder diagnosis; they did not measure baseline cortisol. However, if the individuals in the Demers study were non-minor at CRFR1 rs7209436 and rs242924, then their participants would match our results of FAAH AA or AC being riskier for anxiety in individuals who were not carriers of all three minor alleles. But, the sample populations in the two studies were not identical. In our sample, it is important to note that individuals were not clinically anxious, and our anxiety scores represent natural variation in state (current) anxiety in a population; whereas the individuals in the Demers et al. study met DSM-5 criteria for anxiety disorder. However, data suggest that state, trait, and clinical anxiety are related and therefore biomarkers related to one of these measures may be relevant for all of them. For example, state anxiety increases after stress (Ackerl et al., 2002) and this is true for individuals high and low for trait anxiety (Jezova et al., 2004; Roelofs et al., 2007), thus it is plausible that baseline cortisol at the time of state anxiety measure and genotypes related to anxiety and stress responsiveness would be associated with both state and trait anxiety. Moreover, state and trait anxiety are highly correlated (Spelberger, 1983) and individuals with higher levels of anxiety sensitivity are more at-risk for clinical anxiety disorders (Naragon-Gainey, 2010). Nonetheless, it is possible that biomarkers of state anxiety and those for trait or pathological anxiety differ.

In conclusion, in our study population, the minor alleles of CRFR1 and the AA or AC genotype of FAAH are related to reduced anxiety, but surprisingly, related to higher, and not lower, baseline cortisol. Additionally, the FAAH CC genotype is only confers risk for anxiety in individuals who are also carriers of the CRFR1 minor combination. But, FAAH genotype was not independently associated with anxiety, and baseline cortisol was negatively associated with anxiety score. These results may help to interpret how genetic differences in the HPA axis and EC system impact stress-related disorders.

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Appendix A. Supplementary data

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