



Published in final edited form as:

Biol Psychol. 2020 May ; 153: 107888. doi:10.1016/j.biopsycho.2020.107888.

The interactive effect of family conflict history and physiological reactivity on different forms of aggression in young women

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Abstract

Evidence indicates that patterns of biological reactivity underlie different forms of aggression, but greater precision is needed in research targeting biopsychosocial processes that underlie such differences. This study investigated how sympathetic and parasympathetic nervous system (SNS and PNS) responses to social stress were associated with multiple forms of aggression in an ethnically-diverse sample of young adult females; it further examined whether early life exposure to family conflict moderated these relationships. In the context of high levels of family conflict history, greater SNS activation during a social conflict task was associated with more direct proactive aggression and increasing RSA was associated with more direct reactive aggression. Greater SNS activation during the task was associated with more direct reactive aggression regardless of family conflict history. Our findings affirm the need to capture the contributions of multiple physiological systems simultaneously and the importance of considering family history in the study of aggression.

Keywords

Aggression; Females; Family conflict; Adversity; Stress; Parasympathetic; Sympathetic

1. Introduction

Young adults' behavioral patterns forecast the quality of subsequent interactions with long-term partners and offspring, making this developmental period a critical time for interrupting cycles of aggression. A significant body of literature suggests that patterns of biological reactivity may underlie different forms of aggression, but greater precision is needed in research targeting the specific dynamic biopsychosocial processes that underlie such differences. Notably, studies on the psychophysiology of aggression have historically focused on males and have largely ignored the role of young adults' early family environment, despite the strong influence that childhood experiences exert on long-term

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Declaration of Competing Interest

The authors have no conflicts of interest to report.

behavior problems. To address these gaps in the literature, the present study examined the associations between autonomic reactivity in response to social conflict and different forms of aggression in an ethnically-diverse sample of young adult females. It additionally sought to determine whether these associations are moderated by exposure to family conflict during childhood. In addition to shedding light on biological processes implicated in female aggression, this work has the potential to lead to more specificity in violence prevention research relevant to a developmental period largely overlooked in prevention science.

1.1. Psychophysiological stress reactivity and aggression

Multiple theories recognize that physiological arousal is a key component of aggression (Anderson & Bushman, 2002; Goozen, Fairchild, & Harold, 2007) and that the interaction of psychosocial and biological mechanisms underlying aggressive and violent behavior must be more fully understood (Murray-Close, 2013). Most studies on the physiology of aggression have focused on the autonomic nervous system (ANS), which is composed of the sympathetic (SNS) and parasympathetic (PNS) nervous systems (Murray-Close & Rellini, 2012; Patrick & Verona, 2007). The SNS is responsible for supporting bodily functions during episodes of stress while the PNS is responsible for supporting bodily functions of rest; as such, each are regarded as having opposing effects on the body. Engagement of the SNS during stress underlies the “fight or flight” somatic response, resulting in glucose increases, elevation in heart rate and increased blood pressure. To facilitate this reactivity, withdrawal of the PNS occurs. Stress-induced change in PNS activity is determined by measuring the influence of the tenth cranial nerve, the vagus, on the heart. PNS activation (whereby the PNS is engaged rather than withdrawn during stress) is often referred to as vagal augmentation, or increased influence of the vagus, and vagal withdrawal describes decreased influence (Mendes, 2009).

The body of work on ANS reactivity and aggression has produced inconsistent and often contradictory findings. For example, a number of studies have found that aggression in adulthood is associated with greater ANS reactivity to stress as indicated by elevated sympathetic reactivity (Beauchaine, Gatzke-Kopp, & Mead, 2007; Lorber, 2004; Murray-Close & Crick, 2007). Conversely, other investigations have reported associations between aggression and lower ANS reactivity, as indicated by blunted sympathetic reactivity (Ortiz & Raine, 2004; Posthumus, Böcker, Raaijmakers, Van Engeland, & Matthys, 2009) or blunted withdrawal of the parasympathetic system (i.e., vagal augmentation; Calkins, Graziano, & Keane, 2007; Katz, 2007; Obradovi, Bush, Stamperdahl, Adler, & Boyce, 2010).

This inconsistency in the field is likely due to a number of issues. First, research in this area has tended to rely on non-specific measures of isolated parts of the human stress response system. As an example, an impressive body of research documents negative associations between resting heart rate or heart rate reactivity and aggression or disorders associated with aggression (i.e. conduct disorder, oppositional defiant disorder, antisocial personality disorder) (Lorber, 2004; Ortiz & Raine, 2004). Heart rate reflects both SNS and PNS influences on the heart; the relative contributions of the two systems cannot be disentangled from this measure. A more complete understanding of how the ANS is related to aggression

may be obtained by effectively measuring the two branches of the ANS independently and then examining their associations with aggression separately and in light of the other.

Second, much of the research on the psychophysiology of aggression has lacked specificity in aggression measurement. Several critical distinctions have been suggested regarding different forms of aggression. *Direct aggression* involves both overt physical attacks and verbal assaults of the intended victim, although many studies focus on *physical aggression*, specifically. Conversely, *indirect aggression*, which has also been referred to as *relational aggression*, involves harm delivered circuitously and/or through relational means such as social exclusion or spreading rumors (Crick, Casas, & Mosher, 1997). While there may be differences in the neurobiological underpinnings of direct and indirect aggression, research exploring these distinctions is fairly limited (Murray-Close & Crick, 2007; Murray-Close, Han, Cicchetti, Crick, & Rogosch, 2008; Sijtsema, Shoulberg, & Murray-Close, 2011). Illustratively, there is some evidence that increased ANS reactivity (e.g., increased systolic blood pressure reactivity or heart rate reactivity) is associated with indirect aggression in girls, whereas decreased ANS reactivity (e.g., blunted heart rate reactivity and blunted parasympathetic withdrawal) may be associated with direct aggression in boys and girls, respectively (Murray-Close & Crick, 2007; Sijtsema et al., 2011).

Further, within direct and indirect aggression, theoretical and empirical evidence suggests two unique subtypes: proactive and reactive aggression (Little et al., 2003; Poulin & Boivin, 2000; Stanford et al., 2003). *Proactive aggression* includes behaviors perpetrated for the purpose of gaining resources or control over others (and may or may not occur in response to specific provocation); conversely, *reactive aggression* involves behavioral responses that arise following a perceived threat or provocation (Teten et al., 2011). Although proactive and reactive aggression can co-occur, they serve different functions and are associated with unique patterns of physiological reactivity (Babcock, Tharp, Sharp, Heppner, & Stanford, 2014; Bobadilla, Wampler, & Taylor, 2012; Hubbard et al., 2002; Murray-Close & Rellini, 2012). A recent investigation of the biological underpinnings of reactive and proactive indirect aggression offers the most comprehensive and compelling evidence in young adults to date (Murray-Close, Holterman, Breslend, & Sullivan, 2017). Utilizing a standardized social stress task, they found that blunted withdrawal of the PNS during stress (i.e., vagal augmentation) predicted *proactive relational aggression*, and greater SNS reactivity (changes in skin conductance) predicted *reactive relational aggression*. They also found that vagal augmentation was most likely to predict any form of aggression if the individual also exhibited low skin conductance reactivity.

Murray-Close et al. (2017) made a significant contribution to the field owing to their examination of different forms of stress (cognitive and social) and different forms of aggression, as well as the testing of interactions between multiple arms of the ANS. The current investigation furthers our understanding by employing an ecologically-valid stress task that involves social conflict (thus making it more relevant to aggression and frustration), and drawing upon a more ethnically diverse sample. Latina and African-American women may be three times as likely as White women to experience victimization and two times more likely to engage in aggression toward intimate partners (Field & Caetano, 2005); however, very limited empirical research has focused on aggression in women of color,

particularly during the period of young adulthood (Goldstein, 2011; Rivera-Maestre, 2014). In addition, in the current work, we consider a factor known to exert a critical influence on the development of aggressive behavior: qualities of the early childhood family environment.

1.2. The role of the childhood family environment

Theories of diathesis-stress and differential susceptibility argue that connections between biology and behavior cannot be understood without considering environmental context early in life (Ellis, Boyce, Belsky, Bakermans-Kranenburg, & van Ijzendoorn, 2011); a substantive body of research supports this claim (see review in Bush & Boyce, 2014). These models posit that complex interactions between stress reactivity and social experiences early in life underlie the emergence and maintenance of psychopathology. As outlined above, the majority of studies that have examined the associations between physiological stress reactivity and aggression have not assessed individuals' history of trauma or quality of the childhood family environment (Bohnke, Bertsch, Kruk, & Naumann, 2010; Gordis, Granger, Susman, & Trickett, 2006; Lorber, 2004; Patrick & Verona, 2007; Verona & Kilmer, 2007; Verona & Sullivan, 2008), though there are a few exceptions (El-Sheikh, Hinnant, & Erath, 2011; Obradovi , Bush, & Boyce, 2011; Saxbe, Margolin, Spies Shapiro, & Baucom, 2012).

Despite the strong theoretical and empirical foundation supporting biopsychosocial models of disordered behavior and extensive evidence that individuals raised in aggressive households have a greater risk of developing aggressive behavior patterns later in life (Cui, Durtschi, Donnellan, Lorenz, & Conger, 2010; Herrenkohl & Herrenkohl, 2007), we could identify only a single study that tested a biopsychosocial model of aggression in young adults, specifically. It was found that vagal withdrawal (i.e. PNS reactivity) was positively associated with reactive aggression and negatively associated with proactive aggression in conditions of "low social adversity" (e.g., less poverty, lack of parental incarceration, uncrowded home, smaller family) (Zhang & Gao, 2015). This study provides compelling preliminary evidence of important environmental influences on the association between stress reactivity in response to a non-social task and different forms of aggression in an ethnically diverse, yet relatively small ($n = 84$), sample of young adults.

1.3. The current study

The present study extends the existing base of research in a number of important ways. Specifically, the present study 1) utilizes multiple channels of ANS reactivity in reaction to a stressor; 2) employs differentiated measures of aggression (reactive direct, proactive direct, reactive indirect, and proactive indirect); 3) includes an ethnically-diverse female sample; and 4) includes reports of childhood family conflict in order to test moderating effects. Taken together, the current investigation will be the first study to involve a sophisticated test of the associations between physiological reactivity to stress, relevant developmental experiences, and aggressive behavior outcomes in population that is at elevated risk for aggression yet is underrepresented in the literature. Such an investigation may help to resolve the inconsistent findings in the field regarding associations between psychophysiology and aggression by considering direct associations between each branch of the ANS in the context of childhood exposure to family conflict, while controlling for the other branch of the ANS.

Based on prior examinations of the biological underpinnings of delinquency in children from high conflict families (El-Sheikh et al., 2011) and models of differential susceptibility (Bush & Boyce, 2014), we posited that blunted SNS reactivity to stress will be associated with greater proactive aggression and heightened SNS reactivity will be associated with greater reactive aggression among women exposed to higher levels of family conflict earlier in life, while controlling for PNS activity. We also hypothesized that blunted PNS reactivity (vagal augmentation) and greater PNS reactivity (vagal withdrawal) would be associated with greater proactive and reactive aggression, respectively, among women exposed to higher family conflict while controlling for SNS activity. We had no a priori hypotheses regarding the form of aggression (direct vs. indirect), but based on research with children, we tentatively expected that associations might be stronger for indirect aggression compared to direct aggression given that the sample includes only women and indirect aggression frequency and variability might be higher compared to physical aggression in this population.

2. Methods

2.1. Participants

One hundred seventy-eight female young adults attending a state university or community college in a major city in the Western United States participated in the study. Recruitment procedures included: flyers displayed on bulletin boards and metro lines surrounding the two campuses; scripted announcements in university classrooms; and flyer postings to online and in-person university courses. A special effort was made to recruit women of color by posting on Facebook pages of student groups representing Latinx and Asian students and making announcements in Ethnic Studies-focused courses. Due to the nature of the study, which included neurobiological measures of stress reactivity, eligibility criteria included the following: 18–35 years of age, English speaking, body mass index less than 35, without a history of serious medical conditions (particularly heart or respiratory ones), and not taking medications that could interfere with the assessment of stress physiology.

Of the 178 participants who completed the study, nine failed attention check questions and/or did not speak fluent English, and ten participants were missing physiological data due to equipment failure or experimenter error. The final sample for the current analysis included 159 young women (Mean age: 21.6 years, $SD = 3.15$), including 130 undergraduate students without (52 %) and with (30 %) an Associate's Degree and 29 master's students (18 %). The sample was ethnically diverse: 32 % Latina/Hispanic, 27 % Asian/Pacific-Islander, 24 % White, 6% Black, 9% Multi-ethnic, and 3% Other. Of those who reported on parental nativity ($n = 150$), 50 % indicated that neither parent was born in the United States. The majority of women reported earning less than \$10,000 per year (77.4 %).

2.2. Procedures

All procedures were approved by the university Institutional Review Board. Individuals who were interested in participating in the study were directed to an online registration form that included the eligibility criteria. If eligible, participants were instructed 24 h prior to their appointment to refrain from vigorous exercise, smoking, drinking alcohol, coffee/energy

drinks, consuming food or cold medicine for at least 2 h before their scheduled lab visit. Appointments were either at 1 pm or 4 pm on a weekday and lasted approximately 2 h. Following the informed consent process, height and weight measurements were taken to calculate body mass index. Participants were then attached to physiological equipment. Following a baseline resting period, the participants completed two conflict discussion role play tasks and rated their mood and cognitive appraisals prior to and following the tasks. After detachment from the physiological sensors, participants completed a battery of questionnaires on a computer located in a room separate from where the stress tasks occurred. Participants were compensated with \$25 in the form of an Amazon gift card or extra credit for a psychology course.

2.3. Measures

2.3.1. Self-report questionnaires

2.3.1.1. Direct aggression.: Direct aggression was measured using the Reactive-Proactive Questionnaire (RPQ; Raine et al., 2006), with slight word modifications to ensure it was appropriate for emerging adults. The RPQ includes a 12-item reactive aggression subscale (e.g., “Yelled at others when they have annoyed you.”) and an 11-item proactive aggression subscale (e.g., “Had fights with others to show who was in control.”). Questions were rated on a scale of 0 (*never*), 1 (*sometimes*), or 2 (*often*) and items on each of the two subscales were averaged, with higher scores reflecting more direct reactive or proactive aggression. Given that the RPQ was originally developed for child populations and measurement models have not been published for normative young adult populations, a confirmatory factor analysis (CFA) was conducted. Results showed that 4 items on the proactive subscale loaded poorly ($<.30$). These items were removed. Final analyses utilized the full 12-item reactive aggression scale ($\alpha = .81$) and the reduced 7-item proactive aggression scale ($\alpha = .59$).

2.3.1.2. Indirect aggression.: Indirect aggression was assessed using the peer-directed aggression subscales of the Self-Report of Aggression and Social Behavioral Measure (Murray-Close, Ostrov, Nelson, Crick, & Coccaro, 2010). The original measure included 13 items assessing peer-directed proactive and reactive indirect aggression; however, the current study utilized the nine items employed in a previous study (Murray Close et al., 2010; $\alpha = .81$). Four items assess reactive indirect aggression (e.g., “When I am not invited to do something with a group of people, I will exclude those people from future activities.”) and five items assess proactive indirect aggression (e.g., “My friends know that I will think less of them if they do not do what I want them to.”). Items were rated on a scale from 0 (*never*) to 4 (*very often*) and averaged for each subscale, with higher scores reflecting more indirect aggression. Internal consistencies for the reactive and proactive subscales in the current study were .78 and .65, respectively.

2.3.1.3. Family conflict in childhood.: The Risky Family Environment questionnaire is a 13-item self-report measure that assesses aspects of the childhood family environment, such as chaos, predictability, warmth, and conflict (Taylor, Lerner, Sage, Lehman, & Seeman, 2004). Participants indicated how true each statement was for them on a scale from 1 (*not at all*) to 5 (*very often*). For the current study, only the 7 conflict items were used (e.g., “How often did a parent or other adult in the household push, grab, shove, or slap you?”; $\alpha = .81$).

2.4. Social conflict paradigm

2.4.1. Role play tasks—Participants completed two 5-minute role play tasks (Larkin, Semenchuk, Frazer, Suchday, & Taylor, 1998), the order of which was counterbalanced across the sample. For each task, a trained confederate entered the room and sat facing directly across from the participant approximately 30 in. away. All confederates were instructed to display a neutral facial expression, maintain flat affect, make direct eye contact, and not engage with the participant outside of their assigned lines. In the *Messy Roommate* roleplay, participants interacted with a female confederate in a role play scenario in which participants were instructed to act out the following: “Your roommate is a slob and the apartment is a mess. You always do your share. You ask her to do the dishes because you have friends coming over. You get back home and the place is worse than when you left it. Your goal is to get your roommate to agree to clean up the apartment.” In the *Noisy Neighbor* role play task, participants interacted with a male confederate in a role play scenario in which participants were instructed to act out the following: “You are trying to study for an important exam. You really need to do well on this exam, but you can’t concentrate because your neighbor is playing his music too loud. You decide to ask him to turn down his music so you can study.”

2.4.2. Mood and cognitive appraisals—Participants’ negative mood and negative appraisals of the tasks were assessed prior to and following the role play tasks. The Positive and Negative Affect Schedule is a 20-item self-report measure that assesses current positive and negative affect (PANAS; Watson, Clark, & Tellegen, 1988). Participants indicated how much they currently felt each state on a scale from 1 (*not at all*) to 5 (*extremely*). Only the 10-item Negative Subscale was used in the current study ($\alpha = .83$). The Stress Appraisal Measure is a 28-item self-report measure that assesses an individual’s appraisal of a specific stressful situation (Peacock & Wong, 1990). Participants indicated how they felt about the situation on a scale from 1 (*not at all*) to 5 (*extremely*). Only the 4-item Threat Subscale was used in the current study ($\alpha = .70$). These measures served as a manipulation check for the social stress tasks.

2.5. Physiological measurements

Autonomic Nervous System (ANS) stress reactivity was recorded using electrocardiography (ECG) and impedance cardiography (ICG) with BIOPAC (Biopac MP150, Data Acquisition System, Biopac Systems, Inc., www.biopac.com). The experimenter applied two strips of impedance tape around the circumference of participants’ necks with its mylar 3 cm apart from each other and parallel to the ground. In a similar fashion two other strips were applied around the torso, right below the bra line. Two ECG electrodes were then placed, one on the right breastbone and the other under the left rib cage, slightly above the belly button. Data were sampled at 1000 Hz.

There were a total of four relevant consecutive recording sessions: Baseline, Social Stress task 1, Social Stress task 2 and Recovery, each five minutes long for a total of 20 min of recording per participant. Data was cleaned using Mindware HRV and IMP v3.0.15 by visually inspecting each 1-minute segment for artifacts and edited when needed. Segments in which more than 10 % of the data were missing were excluded from analyses. Minute

segments were averaged together for each recording session to create a single mean score per session.

We focused specifically on Pre-Ejection Period (PEP; a time-based measure that is determined as the time from the left ventricle contracting to the opening of the aortic valve) and Respiratory Sinus Arrhythmia (RSA; a type of heart rate variability in which spectral analysis is used to derive the high frequency component of the interbeat Interval cycle (0.12 to 0.40 Hz)) because they have been known to assess activation of the sympathetic and parasympathetic systems, respectively. Studies using pharmacological blockades have shown that PEP measures cardiac sympathetic activation without the influence of the vagus nerve (Berntson et al., 1994), and RSA is considered mainly vagally mediated, reflecting cardiac parasympathetic activation without sympathetic input (Mendes, 2009). For PEP, a smaller reactivity score means greater activation of SNS, whereas for RSA, a smaller reactivity score means less activation of PNS, both of which are interpreted as higher stress reactivity.

Paired t-tests contrasting average baseline PEP and average PEP during the first task indicated that PEP values decreased significantly from baseline ($M = 105.54$, $SD = 10.92$) to the end of the first task ($M = 90.87$, $SD = 15.36$), $t(157) = 15.65$, $p < .001$, and from baseline to the end of the second task ($M = 93.30$, $SD = 14.06$), $t(156) = 13.82$, $p < .001$. RSA values also decreased from baseline ($M = 6.56$, $SD = 0.97$) to the first task ($M = 6.45$, $SD = 0.90$), although statistical significance was marginal, $t(157) = 1.74$, $p = .09$. There was no change in RSA values from baseline to the second task, ($M = 6.55$, $SD = 0.87$), $t(156) = .12$, $p = .91$. PEP change from baseline to the first task was statistically significantly greater than PEP change from baseline to the second task ($M^{diff} = -2.44$, $t = -5.13$, $p < .0001$). Additionally, RSA change from baseline to the first task was also greater than RSA change from baseline to the second task ($M^{diff} = -0.11$, $t = -3.26$, $p = .001$). We interpret these differences as evidence of habituation to the second task. Thus, reactivity scores to be used in the preliminary and primary analyses were calculated by subtracting average PEP and RSA at baseline from average PEP and RSA during the first task, respectively.

2.6. Data analysis

Study variables were inspected for non-normality and outliers. Inspection of aggression and reactivity scores revealed a number of outliers. One case had a direct proactive aggression value above 3*Interquartile Range (IQR) and was winsorized to the next highest value. Two cases had indirect proactive aggression values more than 3*IQR; these values were winsorized to the next highest value. Four individuals had PEP reactivity scores that were 3*IQR and three individuals had RSA reactivity scores that were 3*IQR. These high levels could not be explained by these participants' body mass index, behavior during the task, or daily use of caffeine, marijuana, or alcohol; therefore, these cases were winsorized to the next highest value and included in the analyses.

Correlational analyses were conducted to examine bivariate relations between the primary study variables and potential covariates, including age, income, and race/ethnicity. Given the ethnic diversity of the sample, we also explored whether the primary variables of interest differed across different ethnic groups. Primary analyses were conducted within a structural equation modeling (SEM) framework in Mplus v.8 using maximum likelihood estimation

with robust standard errors, which is appropriate for non-normally distributed data. The SEM framework allowed for the simultaneous estimation of relations between the independent variables (i.e., PEP reactivity, RSA reactivity, and childhood family conflict) and the four different types of aggression: direct reactive, direct proactive, indirect reactive, and indirect proactive aggression. The three independent variables were mean-centered and interaction terms were created by taking the products of PEP reactivity \times family conflict and RSA reactivity \times family conflict. Specificity analyses were conducted to inspect the influence of a single participant who scored substantially higher on all aggression measures. There was a significant change in multiple coefficients when this participant was excluded, suggesting that this case exerted undue influence on the associations; therefore, all analyses were conducted excluding this one case.

3. Results

3.1. Preliminary analyses

A manipulation check of the social stress tasks was conducted by examining changes in self-reported mood, self-reported negative cognitive appraisals, and autonomic reactivity to each of the tasks. Paired t-tests indicated a significant increase in negative mood from pre-task ($M = 13.63$, $SD = 4.67$) to post-tasks ($M = 18.30$, $SD = 6.51$), $t(157) = -10.92$, $p < .001$ and a significant increase in threat appraisal from pre-task ($M = 1.77$, $SD = 0.62$) to post-tasks ($M = 2.32$, $SD = 0.88$), $t(156) = -8.59$, $p < .001$, providing evidence that participants subjectively experienced the tasks as stressful.

Descriptives and zero-order correlations among the variables of interest are displayed in Table 1. Mean levels of aggression were quite low. In addition, young women endorsed family conflict as happening “rarely” on average; however, 30 % of women reported that family conflict occurred “sometimes” or “often”. Family conflict was associated with greater direct reactive aggression but showed no statistically significant correlation with direct proactive aggression or either form of indirect aggression. Family conflict was also correlated with greater RSA reactivity (i.e., more parasympathetic withdrawal). Income and age were not statistically significantly correlated with any of the primary variables ($ps > .10$, not shown). In addition to the expected positive associations among all aggression indicators, there was also a small but statistically significant positive correlation between PEP reactivity and RSA reactivity. The small to moderate correlations between the different types of direct and indirect aggression indicate the value of examining these as separate outcomes.

To explore whether the primary variables of interest differed across ethnicity, four groups were created based on previous research showing differing levels of aggression in White, Asian, and Latina populations. Due to the low proportion of African-American women ($n = 9$) and other race/ethnicities (e.g., American Indian/Native American/Indigenous) in the current sample, they were collapsed into a fourth “other” category. History of family conflict did not differ across the groups ($p = .84$). In addition, regardless of type or form, aggression did not differ across the four groups ($ps .12-.92$). Finally, although PEP reactivity also did not differ across race/ethnicity ($p = .21$), RSA reactivity was significantly greater (more

parasympathetic withdrawal indicated by larger decreases in RSA values) for women who identified as Latina compared to White women ($M^{\text{diff}} = -.453, p = .04$).

3.2. Primary analyses

Results are displayed in Table 2, organized by type of aggression.

3.2.1. Direct reactive aggression—Greater RSA augmentation (less parasympathetic withdrawal), greater PEP reactivity, and greater family conflict all predicted greater direct reactive aggression. However, these associations must be qualified given the statistically significant interaction between RSA reactivity and family conflict ($p = .038$; Table 2). Simple slopes at $+1 SD$ and $-1 SD$ were estimated and tested following guidelines by Aiken and West (1991) using Mplus and plotted using the PROCESS Macro in SPSS. As shown in Fig. 1, at high levels of family conflict ($+1 SD$), greater RSA augmentation (i.e., less parasympathetic withdrawal, indicated by increased RSA) was associated with higher endorsement of direct reactive aggression, $\beta = .23, SE = .11, p = .03$. At low levels of family conflict ($-1 SD$), RSA reactivity was not associated with direct reactive aggression ($p = .55$).

3.2.2. Direct proactive aggression—The interaction between PEP reactivity and family conflict was statistically significant ($p = .007$; Table 2). Simple slopes at $+1 SD$ and $-1 SD$ were estimated and tested following guidelines by Aiken and West (1991) using Mplus and plotted using the PROCESS Macro in SPSS. Neither simple slope reached conventional levels of statistical significance ($ps > .05$); however, this is not unusual in the case of a full cross-over interaction like the one observed here (Baron & Kenny, 1986). As shown in Fig. 2, at high levels of family conflict ($+1 SD$), greater PEP reactivity (i.e., more SNS activation, indicated by decreased PEP) was marginally associated with higher endorsement of direct proactive aggression, $\beta = -.21, SE = .11, p = .06$. At low levels of family conflict ($-1 SD$), less PEP reactivity was associated with higher endorsement of direct proactive aggression but the effect was not significant, $\beta = .16, SE = .10, p = .10$.

3.2.3. Indirect reactive aggression—Neither interaction term was statistically significant, nor were there any main effects of RSA reactivity or family conflict history on indirect reactive aggression. There was a negative association between PEP reactivity and indirect reactive aggression, but this estimate did not reach statistical significance ($p = .07$; Table 2).

3.2.4. Indirect proactive aggression—The interaction between PEP reactivity and family conflict was marginally significant for indirect proactive aggression ($p = .055$; Table 2). Although the estimate did not reach statistical significance, simple slopes were tested to explore whether the pattern of the interaction mirrored that found for the direct form of proactive aggression. Greater PEP reactivity (i.e., more SNS activation) was associated with higher endorsement of indirect proactive aggression, $\beta = -.24, SE = .10, p = .019$. At low levels of family conflict ($-1 SD$), the association between PEP reactivity and indirect proactive aggression was not statistically significant, $\beta = .13, SE = .11, p = .24$.

4. Discussion

Research has produced inconsistent and often contrary findings, with both high or low ANS reactivity shown to increase the risk of aggressive behavior. Although childhood exposure to family conflict is known to predict aggression later in life, few investigations have examined how family history of aggression and biological processes work together in a dynamic fashion to increase the risk of aggressive behavior, particularly among ethnically-diverse female populations. To address this gap, the current study tested whether childhood family conflict exposure moderated associations between physiological responses to social conflict and different forms of aggression (direct and indirect, reactive and proactive) in an ethnically-diverse sample of young adult females. In doing so, the investigation represents a sophisticated test of the associations between physiological reactivity to stress, relevant developmental experiences, and aggressive behavior outcomes in population that may be at elevated risk for aggression yet is underrepresented in the literature.

First, it was expected that, among women exposed to higher levels of family conflict earlier in life, blunted SNS reactivity to stress would be associated with greater proactive aggression, whereas heightened SNS reactivity would be associated with greater reactive aggression, while controlling for PNS reactivity. Family conflict moderated the association between SNS reactivity and proactive aggression only. Contrary to expectations, heightened SNS reactivity was associated with higher levels of direct proactive aggression among women exposed to high levels of family conflict. This finding is consistent with recent work showing that heightened baseline SNS arousal was associated with greater proactive aggression in young adult women (e.g., Armstrong et al., 2019), but suggests that family conflict exposure represents a critical contextual variable. The role of family conflict in the psychophysiology of proactive aggression is in line with differential susceptibility models of psychological problems, whereby individuals with heightened sensitivity to context are expected to develop particularly problematic behaviors under conditions of adversity. The lack of a correlation between SNS reactivity and family conflict history suggests that heightened sympathetic reactivity did not develop as a result of exposure to conflict. However, given the cross-sectional nature of the study, it is not possible to conclude that young women's pattern of physiological responses were also evident earlier in life.

The association between heightened SNS reactivity and greater proactive aggression for women exposed to family conflict contrasts with previous findings of the opposite association among young women who experienced sexual abuse (Murray-Close & Rellini, 2012). Although sexual abuse may exert a unique effect compared to exposure to family conflict, a number of differences between the two investigations may also account for the divergence in findings, including but not limited to the characteristics of the samples (the present study included an extremely diverse and larger sample of women), the measure of SNS (in the previous study, heart rate reactivity was used as an indicator of ANS response, which captures both sympathetic and parasympathetic influences) and the use of different stress tasks (the previous study employed a social stress interview). The use of an ecologically-valid social stress task in the current study is a unique strength given that stress reactivity to social conflict may be especially salient for those who have been exposed to family conflict.

The current study adds to the growing body of research implicating particular physiological patterns in the expression of different types of aggression, regardless of family history of aggression. It revealed direct associations between each branch of the ANS and multiple measures of aggression while controlling for the other branch of the ANS. This approach contrasts with the common practice of examining biological processes in isolation and is consistent with how these systems actually function in co-ordination within the individual. Consistent with previous research (Hubbard et al., 2002; Murray-Close & Rellini, 2012), heightened SNS reactivity was related to significantly greater direct reactive and marginally indirect reactive aggression, suggesting that women who endorsed the use of reactive aggression became more physiologically activated, particularly in terms of the SNS fight-or-flight response, during social conflict. Reactive aggression may be a behavioral product of the high levels of physiological stress arousal these women experience when faced with perceived antagonism from others. The current study cannot elucidate causal relations, however. It may also be that a history of responding to others with reactive aggression has led to increased levels of physiological activation in response to conflict for these women. Future research employing a longitudinal design could help unpack potential causal pathways.

The association between RSA reactivity and aggression was confined to the reactive form of direct aggression only, was in the opposite direction of expectations, and was strongest for women who reported high levels of family conflict. Lower RSA reactivity or vagal augmentation (indicative of a blunted stress response whereby the PNS does not withdraw as expected) was associated with endorsement of greater direct reactive aggression, *except* for women who reported low levels of family conflict. This finding contrasts with preliminary evidence for an association between higher RSA reactivity (i.e., greater vagal withdrawal) and reactive aggression among young adults who reported experiencing low levels of social adversity (Zhang & Gao, 2015). Because we included both branches of the ANS in the same model we see that PNS activation was accompanied by greater SNS activation (i.e., greater PEP reactivity), which is a state referred to as ANS coactivation (Berntson, Cacioppo, & Quigley, 1991). As described by Murray-Close et al. (2017), both heightened SNS responses to stress and heightened PNS responses (vagal augmentation) to stress are associated with strong or poorly regulated experiences of negative emotion, which may then lead to aggression. Our work indicates that ANS coactivation may be a physiological marker for direct reactive aggression in ethnically-diverse young adult women, especially those exposed to high family conflict.

The present findings must be considered in light of study limitations. Recent research has revealed complicated associations between aggression or aggression-related disorders and interactions between the PNS and SNS. For example, young adults who exhibit low sympathetic reactivity combined heightened parasympathetic reactivity (vagal withdrawal) show greater levels of relational or indirect aggression (Murray-Close et al., 2017). The sample size of the present study did not have sufficient power to test a three-way interaction; future research involving significantly larger samples of participants should consider testing whether history of family conflict or childhood maltreatment influences multi-system stress profiles associated with different forms of aggression. Additionally, although well-validated measures of direct and indirect measures of aggression were used in the current study, the

internal consistency for the measures of proactive forms of aggression were relatively low. Notably, however, the interaction between family conflict and PEP reactivity was present for the two different measures of proactive aggression, adding confidence to the finding. The lower internal consistency observed in the proactive aggression measures may be a result of the lower endorsement of proactive aggression in this population. In fact, overall levels of aggression regardless of form or type appeared to be relatively lower in these women compared to other studies of female aggression (e.g., Dini & Wertag, 2018; Morel, Haden, Meehan, & Papouchis, 2018). Thus, our findings indicate the SNS responses and a history of family conflict are important factors for understanding relatively low, non-clinical levels of aggression.

The current study also has a number of strengths. It advances the field by focusing on a developmental period often overlooked in investigations of aggression. Past research on aggression (as well as prevention/intervention programs) has focused predominately on children and adolescents (Crick & Dodge, 1996) (or on adults with psychiatric disorder; Howells, Daffern, & Day, 2008). However, the majority of at-risk youth do not have access to effective interventions (Mihalopoulos, Vos, Pirkis, & Carter, 2011), which means these same youth enter subsequent developmental stages without having had any corrective interpersonal experiences. Indeed, young adulthood is the most at-risk age group for exhibiting extreme aggressive behaviors, such as homicide (U.S. Department of Justice, 2009), and interpersonal violence disproportionately occurs in young adulthood compared to other developmental stages (Kim, Laurent, Capaldi, & Feingold, 2008). As such, young adulthood a critical time to interrupt the intergenerational transmission of aggression. Our findings demonstrate the importance of assessing the influences of multiple physiological systems on outcomes simultaneously as well as the need to consider the role of developmental history to more fully understand relationships between physiology and behavior. The identification of biological markers associated with different forms of aggression in diverse young women helps inform the development of prevention programs to break the intergenerational transmission of violent behavior in an under-studied and under-served population.

Acknowledgements

This work was supported by grants from the San Francisco State University (SFSU) Office of Research and Sponsored Programs and the National Institute of Health Building Infrastructure Leading to Diversity (BUILD) Initiative (UL1GM118985).

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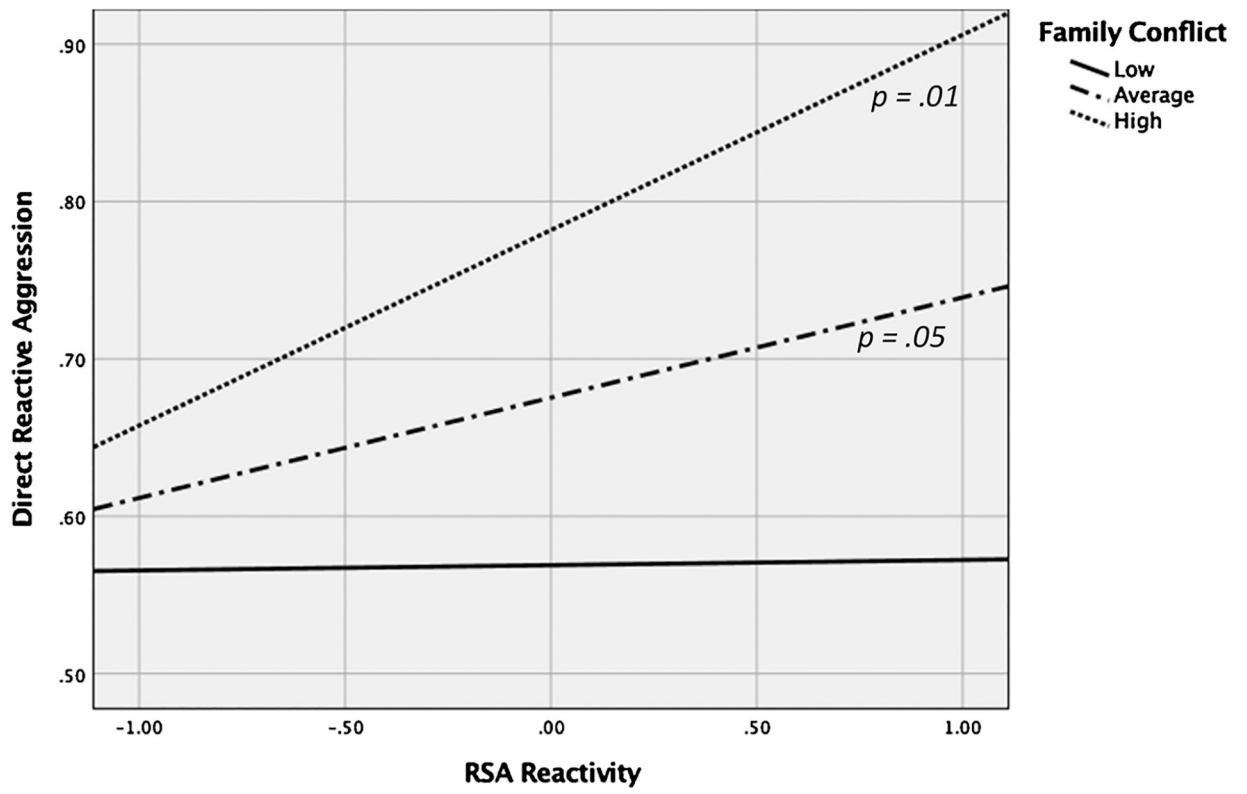


Fig. 1. RSA reactivity predicting direct reactive aggression at +1/-1 SD and mean levels of childhood family conflict.

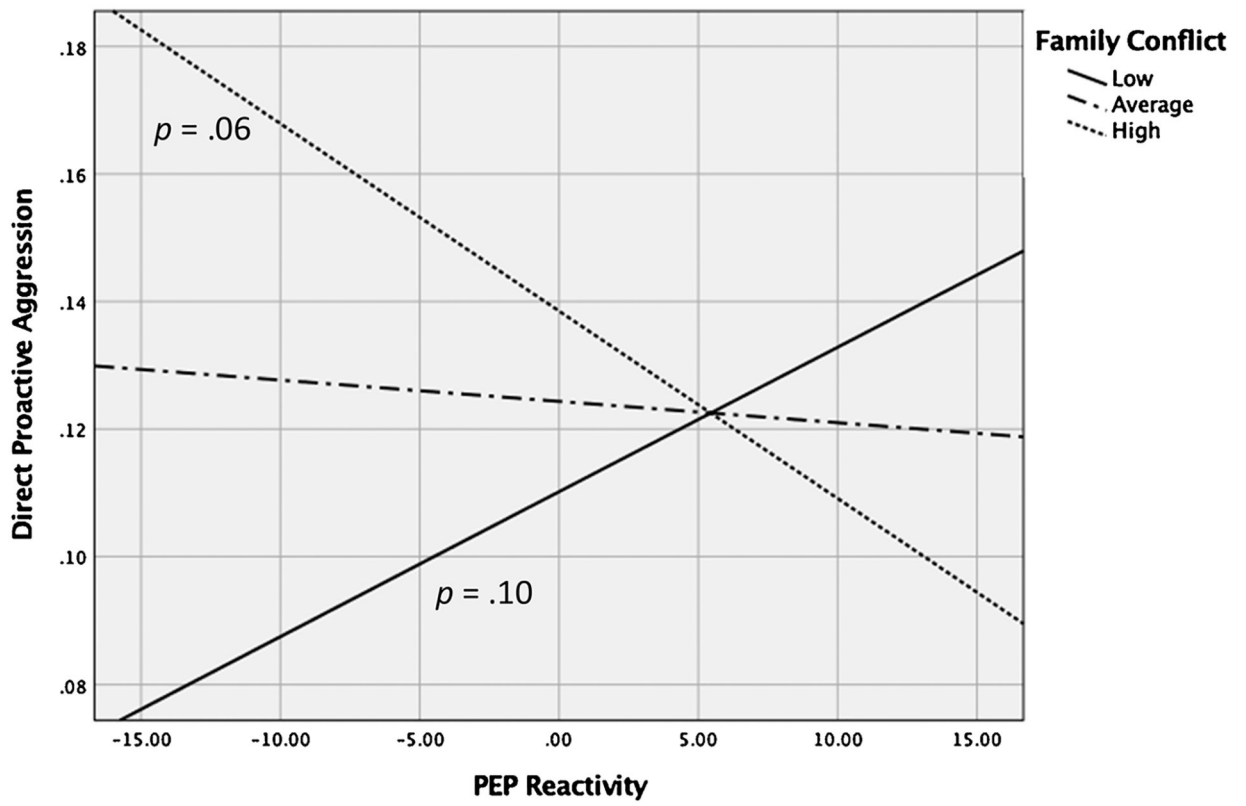


Fig. 2.
PEP reactivity predicting direct proactive aggression at +1/-1 SD and mean levels of childhood family conflict.

Table 1

Kendall's Tau Correlations.

	M (SD)	1	2	3	4	5	6
1. Family Conflict	2.70 (.82)						
2. PEP Reactivity	-14.47 (11.23)	-.03					
3. RSA Reactivity	-.11 (.78)	-.13*	.17**				
4. Direct Reactive	.67 (.34)	.19**	-.08	.02			
5. Direct Proactive	.12 (.16)	.05	-.04	.004	.42**		
6. Indirect Reactive	.52 (.61)	.07	-.04	.02	.36**	.36**	
7. Indirect Proactive	.35 (.42)	-.02	-.03	.08	.27**	.32**	.44**

Note: Negative RSA and PEP reactivity values = more reactivity so the negative correlation between PEP reactivity and reactive aggression is interpreted such that more PEP reactivity is associated with more aggression.

**
 $p < .01$.

*
 $p < .05$.

PEP reactivity interacting with family conflict (Conflict*PEP) and RSA reactivity interacting with family conflict (Conflict*RSA): Standardized estimates and standard errors.

Table 2

	Direct Aggression			Indirect Aggression		
	Reactive		Proactive	Reactive		Proactive
	<i>B</i>	<i>SE</i>	<i>B</i>	<i>SE</i>	<i>B</i>	<i>SE</i>
Family Conflict	.309 ^{**}	.071	.089	.084	.095	.075
RSA reactivity	.146 [*]	.076	-.005	.089	.095	.091
PEP reactivity	-.153 [*]	.073	-.027	.071	-.154 [†]	.083
Conflict*PEP	-.124 [†]	.069	-.225 [*]	.083	-.056	.108
Conflict*RSA	.135 [*]	.065	.131	.082	.062	.082

^{**} $p < .001$.

^{*} $p = < .05$.

[†] $p < .10$.