Pediatric Irritability: A Systems Neuroscience Approach

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Irritability, defined as an increased propensity to exhibit increased anger relative to one’s peers, is a common clinical problem in youth. Irritability can be conceptualized as aberrant responses to frustration (where frustration is the emotional response to blocked goal attainment) and/or aberrant ‘approach’ responses to threat. Irritable youth show hyper-reactivity to threat mediated by dysfunction in amygdala, medial prefrontal cortex (mPFC), anterior cingulate cortex (ACC), insula, striatum, and association cortex. Irritable youth also show abnormalities in reward learning, cognitive control, and responses to frustration. These abnormalities are mediated by circuitry that includes the inferior frontal gyrus (iFG), striatum, ACC, and parietal cortex. Effective treatments for irritability are lacking, but pathophysiological research could lead to more precisely targeted interventions.

Irritability in Children

Irritability (see Glossary) can be defined as an increased propensity to exhibit anger relative to one’s peers. Recently, this clinical problem has become the focus of considerable research interest in child psychiatry and clinical neuroscience [1]. This interest stems from recognition of the irritability’s clinical importance of irritability, given that it is one of the most common reasons children present for mental healthcare [2]. Reflecting this, irritability is the primary feature of a new diagnosis in DSM-5, Disruptive Mood Dysregulation Disorder (DMDD) [3]. Moreover, irritability is prominent in other childhood psychiatric illnesses, including oppositional defiant disorder, anxiety disorders, attention deficit hyperactivity disorder (ADHD), post-traumatic stress disorder, conduct disorder, major depressive disorder, bipolar disorder (BD), and autism spectrum disorders.

Youth with DMDD suffer significant impairment and often require multiple clinical interventions (e.g., medication, school placement, and individual and family psychotherapies) to function adequately at home and school [4,5]. However, the efficacy of commonly used treatments for irritability is limited or, in some instances, unknown. Most such treatments are not designed to target irritability specifically and none are based on an understanding of the relevant neurobiology (for a promising avenue that could be an exception, see [6]). Recent work has yielded much information about the presentation, course, and impact of irritability in youth (Box 1). The next challenge is clear: to elucidate the neural mechanisms of irritability in order to guide the development of novel interventions. Pathophysiological studies targeting irritability specifically remain rare, but there is a considerable foundation of work on related phenotypes, such as reactive aggression (Box 2). Within the realm of psychopathology, irritability is a relatively tractable research target because it is an evoked response and, hence, can be modeled in animals and studied in real time during neuroimaging. Since research on the neural mechanisms of irritability is relatively nascent and rapidly evolving, it is important for investigators to specify neuroscientific conceptualizations to guide research, highlight areas where emerging data warrant follow-up, and discuss approaches to developing promising research paradigms.
suggests that irritability reflects dysfunction in the amygdala-hypothalamic-periaqueductal gray (PAG) threat response circuitry, such that approach responses occur in contexts where the normative response would be freezing or flight [7]. Thus, the common co-occurrence of anxiety and irritability, as well as the genetic and longitudinal links between them (Boxes 1 and 2), may represent vacillation between abnormal avoid and approach responses to threat, reflecting disequilibrium in the mediating circuitry that is, in part, genetically mediated. As discussed below, data in irritable youth and related phenotypes demonstrate abnormalities in threat processing specifically, and in social information processing more broadly. Also as discussed below, this theory has been probed using threat stimuli that are relatively simple, such as angry faces, or that involve more complex social interaction paradigms.

The second hypothesis suggests associations between irritability and abnormal reward processing, specifically in the form of aberrant responses to frustrative nonreward (FNR). In a landmark study, Amsel defined FNR as the psychological state induced by the failure to receive a reward that a rodent has been conditioned to expect. Amsel showed that FNR is associated with increased motor activity and aggression [8]. Research has documented FNR responses in nonhuman primates and humans [9–11] (Figure 1). This work is also clinically relevant, since temper outbursts in irritable children often occur in response to frustration. Taken together, these basic and clinical data suggest that pathological temper outbursts in children reflect FNR responses that are abnormal in their intensity, duration, and/or the strength of the provocation.
Box 2. Clinical Phenotypes Relevant to Research on Irritability

In DSM-5, the pediatric diagnostic categories most germane to the study of irritability are DMDD and oppositional defiant disorder (ODD). Research on irritability is also well suited to a dimensional approach. This is because irritability is distributed continuously across the population and because the precise cut-point differentiating typical from atypical irritability is unclear. Moreover, irritability occurs in many diagnoses (anxiety and ADHD), and a continuous approach facilitates quantification of multiple symptoms simultaneously. Most youth with DMDD also meet diagnostic criteria for ADHD, and many outline diagnostic criteria for an anxiety disorder [4,13]. Conversely, youth diagnosed with ADHD or anxiety exhibit high rates of irritability [64,94]. This co-occurrence among irritability, anxiety, and ADHD may reflect the contribution of both dysfunctional threat processing and attentional dyscontrol to the pathophysiology of irritability. This co-occurrence also raises an important question regarding the extent to which irritability manifesting in different diagnostic contexts is mediated by overlapping or separable neural circuitry (see main text).

As noted in Box 1 (main text), irritable youth may exhibit aggressive behavior during a temper outburst. The brain mechanisms of aggression have received considerable research attention, and studies of reactive aggression are particularly germane to irritability. Reactive or affective aggression is an intense, impulsive behavioral expression of anger, usually precipitated by frustration or threat [95]. Thus, reactive aggression can be seen as the most extreme behavioral manifestation of irritability. Another type of aggression has been termed ‘proactive’ or instrumental. This type of aggression is designed to accomplish a goal (e.g., steal money) and is seen in psychopathic individuals and those with callous-unemotional traits. The pathophysiology of reactive and proactive aggression appears to differ (see main text). As clinical syndromes, reactive aggression is more common than proactive aggression and often exists without co-occurring proactive aggression, while proactive aggression almost universally co-occurs with reactive aggression. This complex clinical pattern complicates the application of findings in the aggression literature to questions about the pathophysiology of irritability. However, work differentiating aggression in psychopathic adults versus those with externalizing traits [96], or in youth with disruptive behavior disorders with or without callous-unemotional traits (e.g., [35,44]; see main text), is helpful in differentiating the pathophysiology of reactive from proactive aggression and, thus, in advancing research on irritability.

Importantly, while stated as dissociable hypotheses, the threat and reward formulations of irritability are inextricably intertwined, both conceptually and empirically. In an extension of Amsel’s work, one study demonstrated an interaction between FNR and threat processing, in that frustrated mice exhibited increased aggressive responses in an intruder paradigm [12] (Figure 1). Indeed, based on other rodent research, Gray [13] proposed the ‘fear = frustration’ hypothesis, suggesting that frustration is processed as a threat and responses to both are mediated by the behavioral inhibition system. A similar formulation emerges from research in humans (i.e., in paradigms that model competitive games, being ostracized or treated unfairly has been viewed by investigators as either a frustrating or a threatening experience [14,15]). Consistent with Gray’s formulation, some research [16] suggests that regions mediating threat processing and reactive aggression (i.e., amygdala, PAG, and insula and dorsal ACC) are engaged during blocked reward. By contrast, other studies [17,18] found competitive processes between regions responding to threat of shock and those engaged in decision-making about a monetary reward. However, this work is only beginning. Important questions suggested by these conflicting data include the impact of the features of the threatening or rewarding stimulus (simple versus complex, social versus nonsocial) on the circuitry engaged, as well as whether distinct clinical phenotypes of irritability have a stronger association with reward- versus threat-related dysfunction.

**Social Information Processing in Irritable Youth**

Irritability in youth usually manifests in social situations, suggesting that irritable youth experience such situations differently than do nonirritable youth. Indeed, multiple lines of research indicate that irritable youth have social information-processing deficits.

**Proactive aggression:** aggression designed to accomplish a goal (e.g., theft), generally exhibited by individuals with psychopathy and/or a tendency to respond to emotional stimuli in a callous and unemotional way.

**Reactive aggression:** aggression that occurs as part of an intense, angry emotional response, usually precipitated by frustration or threat.

**Reward:** a stimulus associated with approach behavior.

**Reward learning:** the process by which organisms learn associations between either a stimulus or behavior, on the one hand, and reward or punishment, on the other.

**Threat:** a stimulus associated with avoid behavior.

**Threat bias:** the tendency to orient preferentially and rapidly to a threatening, versus neutral stimulus.

**Tonic irritability:** irritability manifest as chronically angry mood.
Irritability in Rodents and Humans Is Associated with Aberrant Responses to Threat and/or Frustrative Nonreward (FNR)

Figure 1. The figure shows parallel processes that can be observed in rodents and humans. (A) Associations between aberrant responses to threat and irritability in both species. Experimental manipulations that expose an organism to threat can be used to probe this pathway to irritability. (i) When exposed to the threat of a conspecific in a resident intruder paradigm, frustrated mice show increased aggressive behavior. This aggression is greater than that exhibited by nonfrustrated mice (not shown) [12]. (ii) Irritability in youth is associated with aberrant amygdala-prefrontal cortex connectivity when viewing threatening faces [37]; see Figure 2 (main text) for details. (B) Associations between aberrant responses to frustration and irritability in both species. Experimental manipulations that expose an organism to threat can be used to study between-strain or between-subject differences in response to frustration, in the form of irritable behavior. (i) Frustration is induced in a mouse by

(Figure legend continued on the bottom of the next page.)
The most consistent line of research on social information-processing deficits in irritable youth finds perturbed threat processing. One body of research assesses irritable youth’s responses to relatively simple, iconic social signals, particularly angry faces (e.g., Figure 2), whereas other studies use paradigms that model more complex social processes. In the face emotion-processing literature, paradigms vary in the duration of stimulus exposure and in the cognitive processes engaged. One set of studies uses paradigms such as the dot probe task to assess predicted connectivity change.

Figure 2. Amygdala–Prefrontal Cortex (PFC) Connectivity Is Driven by an Interaction between Levels of Anxiety and Levels of Irritability When Youth View a Threatening Face. For details, see [37]. Briefly summarized, fMRI data were obtained while youth [ages 8–17 y, N = 93 with anxiety, disruptive mood dysregulation disorder (DMDD) and/or attention deficit hyperactivity disorder (ADHD), and 22 healthy volunteers] performed an implicit face emotion-processing task (i.e., gender identification) on angry, happy, and fearful faces at 50%, 100%, and 150% intensity. Functional connectivity was examined using a psychophysiological interaction analysis with an amygdala seed. When subjects viewed a 150% angry face, amygdala connectivity was related to levels of irritability, anxiety, and their interaction. As illustrated in the 3D plane, an interaction between irritability (measured on the ARI scale) and anxiety (measured on the SCARED scale) was found for amygdala–medial PFC connectivity. Specifically, decreasing connectivity was associated with increasing levels of both irritability and anxiety, while increasing connectivity was associated with increasing anxiety in the absence of increasing irritability SCARED, Screen for Child Related Anxiety Disorders.

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individual differences in attention-related threat bias (i.e., the tendency to orient preferentially and rapidly to an angry versus neutral face). In these studies, the face stimulus is presented for a relatively brief period (e.g., 17–1250 ms) and serves as a distractor. Dot probe studies in both clinical and community samples of irritable youth find an attention bias toward angry faces [19,20]. Similar findings manifest in youth with anxiety disorders and have given rise to a treatment (attention bias modification training) designed to decrease anxiety by reversing the threat bias [21,22]. Shared deficits in attention orienting provide one possible pathophysiological basis for the cross-sectional, genetic, and longitudinal associations between anxiety and irritability. To date, there are no attention bias modification training trials in irritable youth, and no fMRI studies in irritable youth that use an attention-orienting task. However, studies of the dot probe task in anxious youth most consistently find abnormalities in amygdala-ventromedial PFC connectivity [23–25].

In other face emotion-processing paradigms, participants are asked to process a specified aspect of the stimulus. Typically, the face is shown for approximately 2000–4000 ms and the subject labels either the emotion (explicit face emotion processing) or another facial feature, such as gender (implicit processing). Given the higher-order cognitive processing involved in emotional labeling, explicit processing tasks would be expected to engage association cortex, whereas some evidence suggests that implicit face-processing tasks activate the amygdala more reliably [26]. On explicit processing tasks, irritable youth show behavioral deficits, in that they label face emotions inaccurately [27,28]. They also judge neutral faces as more hostile than do healthy youth, whether the stimulus duration is brief (i.e., 200 ms) [29] or not (i.e., 4000 ms) [30]. Irritable youth show amygdala, ventral visual stream, and association cortex dysfunction during explicit face processing [30–32], although the nature of the amygdala dysfunction (hyperversus hyperactivation) varies across tasks.

As noted above, compared with explicit face-processing tasks, implicit tasks may be more efficient at eliciting between-group differences in amygdala activity because they are less likely to elicit PFC engagement and, hence, possible amygdala inhibition [26]. Reactive aggression can be viewed as the most extreme behavioral manifestation of irritability. Thus, it is interesting that studies of youth recruited for reactive aggression or irritability find amygdala hyperactivation to implicit processing of negatively valenced faces [33–35]. By contrast, subjects in these studies who exhibited proactive aggression showed amygdala hypoactivation on such paradigms [34,35]. This is consistent with the formulation that youth with irritability or reactive aggression have impulsive ‘hot’ emotional responses to threat, whereas those with proactive aggression process threat in a more considered, ‘cold’ fashion. Analyses of fMRI data acquired at rest or during an implicit face-processing task have also found associations between youth irritability and aberrant activity in other regions, including insula, cingulate, and striatum, but such findings bear replication [33,36,37].

An important question is whether the neural mechanisms mediating irritability vary with differences in clinical phenotype, since irritability can occur with or without various other clinical problems. This includes reactive aggression, and pediatric mental disorders that include DMDD, ADHD, anxiety disorders, and pediatric BD. Is the pathophysiology of irritability similar across these clinical contexts? Two studies address this question using face emotion-processing tasks (one implicit [37], one explicit [32]). Data from both studies suggest that the clinical context moderates the neural correlates of irritability, albeit in somewhat different ways. In the study using explicit face emotion labeling [32], diagnosis was the key parameter influencing neural correlates: children with BD and those with DMDD differed in amygdala and ventral visual stream activity while labeling face emotions. However, in the study involving implicit face emotion processing [37] (Figure 2), severity on an anxiety dimension was the key parameter. Specifically, in the latter study, independent of diagnosis
youth with high levels of both irritability and anxiety tended to have low amygdala–medial PFC connectivity when processing angry faces implicitly, whereas those with anxiety alone tended to have high connectivity. These two studies [32,37] differ in many aspects, including task and the diagnoses compared (of note, BD is a particularly distinct phenotype that is highly heritable [38]); clearly, this important question requires further study.

While paradigms that use face stimuli have the advantage of relatively tight experimental control, they lack ecological validity. To increase the latter, investigators have devised fMRI paradigms designed to model reactive aggression. In paradigms such as the Ultimatum Game [39] or Point Subtraction Aggression Paradigm [40], subjects are told that they are playing a computer game with another person, although the actions of the other ‘person’ are programmed by the investigator. These paradigms assess correlates of reactive aggression (again, an extreme behavioral expression of irritability) by quantifying the extent to which a subject engages in uncooperative or retaliatory behavior in response to the other ‘player’ stealing money, making an unfair offer, or rejecting an ambiguous offer.

While few studies apply these paradigms to children with psychopathology, the results that exist are intriguing. Children recruited risk for either irritability or reactive aggression showed an increase in uncooperative or aggressive behaviors toward the other ‘player’, including in response to ambiguous behavior [41–44]. Thus, irritable or aggressive youth have a ‘hostile interpretation bias’, which can be expressed in diverse contexts and studied from multiple scientific perspectives. On simpler paradigms, this bias could lead irritable children to label ambiguous faces as angry, while, on more complex paradigms, this bias could cause irritable youth to interpret ambiguous peer behavior as threatening [29,45]. Studies suggest that aggressive responding on complex paradigms such as the Point Subtraction Aggression Paradigm or Ultimatum Game is associated with aberrant activity in a threat-mediating circuit including the amygdala, PAG, ventromedial PFC, striatum, and insula, along with decreased amygdala–ventromedial PFC connectivity [43,44]. The ventromedial PFC findings are consistent with an extensive literature suggesting that ventromedial PFC lesions in adult- or childhood cause irritability and aggression [46–48].

In sum, youth with irritability and related phenotypes show hyperreactivity to negatively valenced social stimuli. This manifests as biased orienting to angry faces and as a bias toward interpreting ambiguous social signals as more threatening compared with healthy youth. These biases are associated with amygdala dysfunction, most commonly hyperactivation during implicit processing of negatively valenced faces and decreased connectivity with mPFC. In addition, such biases may be associated with abnormal activation in the ACC, insula, striatum, and association cortex.

**Reward Processing in Irritable Youth**

The FNR model posits that animals respond to the omission of an expected reward with increased activity and aggression; this raises the possibility that irritable youth’s outbursts represent exaggerated FNR responses. Reward-learning deficits in irritable youth could decrease their ability to respond appropriately to reward contingencies, increasing both the probability of frustrating experiences and maladaptive responses to them. Consistent with these possibilities, studies suggest that irritable youth have deficits in passive avoidance and reversal learning, and fMRI studies find associated dysfunction in regions that mediate these psychological processes, including ventromedial PFC, striatum, insula, ACC, and IFG [49–53]. Computational modeling suggests that a decreased ability to represent expected value information in some of these regions, including insula, IFG, caudate, and ventromedial PFC, contributes to impaired decision-making in irritable youth [51,52,54].
Beyond the difficulty in representing expected value, deficient reward learning could arise from impairments in other cognitive control functions. These include the ability to withhold a prepotent response, use working memory to deploy behavior flexibly in complex scenarios, or register and change stimulus–response mappings after making an error. In fact, an extensive literature links pediatric aggression to these and other components of cognitive control [55–58]. While these studies generally do not differentiate reactive from proactive aggression, one study did find a specific association between reactive aggression and deficits in sustained attention and set-shifting [59]. Further evidence of impaired cognitive control in irritable youth arises from imaging and electrophysiology studies. For example, prior studies demonstrated an association between inhibitory control deficits in irritable youth and reduced cognitive control capacities. The latter is reflected in the amplitude of event-related potential components, such as N2 and P3 amplitudes [60–62].

Importantly, irritability, cognitive control deficits, and decision-making abnormalities are all common in youth with ADHD. Irritability is elevated in both community and clinic samples of ADHD, with a tenfold increase in one large community sample and a range of 24–50% in clinic samples [63,64]. While the mechanisms that support associations between ADHD and irritability remain unclear, there is relevant research. A recent study that subtyped patients with ADHD (N = 437) using resting-state fMRI, peripheral physiological methods, and clinical outcome detected an irritable subtype that showed weak parasympathetic response to negative emotional stimuli, reduced amygdala–insula connectivity, and relatively poor outcome [65]. Regarding cognitive control deficits in ADHD, working memory impairment is among the most common [66]. Working memory is thought to play an important role in emotion regulation, in part through facilitating cognitive reappraisal and adaptive responding to changing environmental contexts [67]. During an n-back working memory task that included emotional stimuli, youth with ADHD showed abnormal engagement in multiple PFC regions and striatum [68]. In youth with ADHD, working memory deficits may also contribute to suboptimal decision-making in emotional contexts [69]. Indeed, several studies in youth with ADHD have found impulsive decision-making, assessed using delayed discounting tasks; impulsive responding to threat or frustration is characteristic of irritability and reactive aggression. On delayed discounting tasks, youth with ADHD showed a preference for small, immediate rewards over large, delayed rewards [64,70]. Studies have found associations between such impulsive decision-making and both nucleus accumbens-PFC connectivity and amygdala hyperactivity in youth with ADHD [71,72].

As noted above, studies of perturbed threat processing in irritable youth rely on diverse forms of social stimuli, including both simple iconic images and more complex social game paradigms. Similar diversity applies in studies of perturbed reward processing in irritable youth. Thus, some studies use relatively simple paradigms to assess basic reward learning or cognitive control functions in irritable youth, while others use more complex paradigms that model FNR directly. In the latter, frustration is induced by rigged games or unsolvable puzzles that withhold an expected reward, and investigators report associations between irritability and the degree of frustration induced by the task [62,73–75]. These studies typically find associations between irritability and dysfunction in regions mediating executive attention and reward processing. For example, in a study using functional near-infrared spectroscopy imaging in a sample of preschool children, increased lateral PFC activity was associated with increased frustration tolerance [76]. In older youth, studies of frustration during fMRI or magnetoencephalography showed associations between irritability and ACC, striatal, medial PFC, parietal, and amygdala activity [73–75,77] (Figure 3). Somewhat unexpectedly, in two fMRI studies, irritability was associated with decreased amygdala activation during frustration [74,75]. However, the time course of the frustrative response can be prolonged, complicating the interpretation of baseline activity. The design of such paradigms is also
challenging because of potential order effects (frustration cannot be ‘turned off’ immediately), and because the frustration must be potent enough to be effective but not so potent that the child can not tolerate it and/or moves during scanning. In addition, such tasks often include an element of deception, which requires careful ethical consideration and can also be logistically difficult. Thus, frustration tasks are challenging to implement and engage several

Figure 3. Amygdala Deactivation Differs between Chronically Irritable Youth and Healthy Volunteers While Playing a Frustrating Game. Frustration is induced in children by first inducing reward expectation [i.e., children play a game in which they can easily earn a reward (nonfrustration condition)]. This nonfrustration condition is then followed by a frustration condition, during which the game is rigged so that the child no longer receives the reward he or she has come to expect. Physiological measures, such as skin conductance and BOLD fMRI signal, can be measured while the child plays. The data here are from an fMRI study in which youth with severe mood dysregulation (SMD, the research precursor to disruptive mood dysregulation disorder (DMDD)) and healthy youth played a frustrating game. During the frustration condition, youth with SMD showed decreased amygdala activity when receiving frustrating, negative feedback (‘you lose’). For details, see [74].
complex psychological phenomena. Nonetheless, the ability to evoke FNR directly in the scanner is an important tool in the effort to elucidate the neural mechanisms of pediatric irritability.

Ironically, Amsel’s FNR studies are motivating human research currently, but there have been relatively few recent studies extending his seminal work in rodents. This is unfortunate because, arguably, the most effective translational work in psychiatry has been in anxiety disorders and substance abuse [78,79], and these disorders have salient commonalities with irritability. Specifically, these three phenotypes can all be conceptualized as evoked responses, making it feasible to design parallel paradigms in humans and animals that model the relevant psychological processes. In the case of irritability, one important line of research would be to identify mouse strains that are highly reactive to FNR (e.g., in terms of aggression and motor activity; see Outstanding Questions). This would enable more precise mapping of circuits that are activated differentially in irritable animals. The assumption is that interstrain variability will be larger than intrastrain variability, although, should the latter be more prominent, this would provide an excellent opportunity to focus on individual differences, as is done in research on human psychopathology.

Concluding Remarks
Motivated by its clinical importance, researchers have begun to elucidate the neural correlates of pediatric irritability. In irritable youth, researchers have identified abnormalities in both threat and reward processing. Studies consistently show hyper-reactivity toward threat in irritable youth. Specific findings include an attentional bias toward threat, a tendency to view ambiguous faces as angry, and a propensity to interpret a competitor’s ambiguous actions as hostile. These abnormalities are associated with dysfunction in the circuitry mediating threat processing (amygdala, PAG, and ventromedial PFC) as well as other regions, depending on the behavioral deficit.

Irritable youth also show abnormalities on paradigms that model FNR; those that assess more basic aspects of reward processing; and those that probe cognitive control functions that impact reward processing. Such deficits are associated with dysfunction in the well-defined circuitry mediating reward learning, including ventromedial PFC, insula, ACC, striatum, and IFG. Paradigms modeling FNR also evoke deficits in executive attention and parietal dysfunction in irritable youth. Impaired cognitive control, particularly during inhibition, is present in irritable youth; such deficits, as well as impulsive decision-making, are present in youth with ADHD, who are at particularly high risk for irritability.

Of course, since neurally based research on irritability is relatively new, there are many questions to be addressed (see Outstanding Questions). At the broadest level, it is important to determine how threat-based and reward-based pathways to irritability interact both neurally and clinically, and how the neural circuitry mediating irritability varies across clinical contexts. The former question could be addressed by paradigms that examine associations of irritability with threat and reward processing at baseline, when a threatening stimulus is presented in the context of a rewarding stimulus, and vice versa. It is unknown whether, among irritable youth, there are distinguishable clinical subtypes in which dysfunctional reward or threat processing is more prominent; addressing this question is consistent with the new emphasis on personalized approaches to medical diagnosis and treatment [80]. Similarly, neural specificity within the broad phenotype of irritability can be probed by studies examining whether the circuitry mediating irritability varies across clinical context (e.g., in the presence of reactive aggression, across different diagnoses, etc.), as well as by studies examining how irritability and other traits (e.g., anxiety) [37] interact in their impact on brain function.

**Outstanding Questions**

In rodents, is there interstrain and intrastrain variability in the motor activity and aggression stimulated by FNR? What differences in circuitry function mediate such variability?

How does the circuitry mediating irritability differ across clinical contexts (e.g., in the presence or absence of anxiety, reactive aggression, or ADHD)?

How do threat-based and reward-based pathways to irritability interact neurally and clinically? Can one define subtypes of irritable youth based on the mediating mechanism and, if so, what are the treatment implications of such subtyping?

How does dysfunction in threat reward circuitry differ in irritable youth (who frequently also have anxiety disorders) versus those with anxiety alone?

Reactive aggression can be seen as the most extreme behavioral manifestation of irritability. Are the neural mechanisms mediating these two clinically defined phenotypes on a continuum?

Deficits in both inhibition and working memory have been implicated in the pathophysiology of irritability. What cognitive control deficits are most relevant to the pathophysiology of irritability?

Are irritable youth hyper-reactive to nonsocial threat, as well as to social threat?

If the hyper-reactivity of irritable youth to social threat is ameliorated by computer-based implicit training, would this intervention be effective in decreasing irritability?

At baseline, youth with irritability have cognitive control deficits. Are such deficits exacerbated in the context of FNR? Relatedly, what is the precise nature of the attentional dysfunction that irritable youth exhibit when frustrated?

What deficits in the computational mechanisms mediating reward learning are important in irritability?
Beyond these broad research questions, more specific directions for future research are suggested by the preceding review. To advance research on social information processing in youth with irritability or reactive aggression, it is important to integrate work on these two related phenotypes through better behavioral phenotyping in youth recruited for irritability and better assessment of emotional symptoms in youth recruited for reactive aggression. Few studies of youth recruited for irritability use paradigms designed to study individual differences in ‘approach’ behavior in response to threat, or in the mediating circuitry; the use of such paradigms would also bridge work with reactive aggression. To elucidate reward processing dysfunction associated with irritability, researchers could expand the use of computational approaches to define precisely the relevant learning deficits. Finally, there are relatively few studies of cognitive control function in irritable children at baseline and during frustration; the results of such studies could guide the development of novel interventions.

Understanding the pathophysiology of any psychiatric symptom or syndrome is challenging, and this is particularly true in children, where multiple problems typically co-occur. Irritability is an important focus for research because it is common and associated with both current and long-term impairment, and because few effective treatments exist. Thus, there is a public health imperative to define the neural mechanisms mediating irritability and to use this knowledge to guide the development of novel interventions. Fortunately, there are reasons to believe that irritability may be a relatively tractable target for research, and considerable progress has been made over a relatively short period of time. Further work that brings relief to affected children and their families would be most welcome.

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