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Uniting adolescent neuroimaging and treatment research:

Recommendations in pursuit of improved integration

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Highlights:

- Many clinicians find developmental neuroscience data compelling.
- Neuroimaging protocols can feel far removed from clinical application.
- Practical steps are provided to integrate neuroscience with adolescent clinical advances.

Abstract

Many clinicians who provide mental health treatment find developmental neuroscience discoveries to be exciting. However, the utility of these findings often seem far removed from everyday clinical care. Thus, the goal of this article is to offer, in the context of adolescent mental health, a bridge to connect the fields of applied adolescent treatment and developmental neuroscience investigation. Concretely, an overview of the relevance of developmental neuroscience in adolescent direct practice is provided. A rationale is offered for how and why the integration of neuroscience into the study of adolescent treatment response could benefit adolescent treatment outcomes. Finally, a series of practical suggestions is generated for improving integration of basic science and psychotherapy research, to enhance collaborative, interdisciplinary work that ultimately advances treatment response for this important clinical population.

Key words: clinical, adolescents, developmental neuroscience

Uniting adolescent neuroimaging and treatment research:

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From the onset of puberty through age 25, the adolescent brain undergoes profound changes in structure and function [1]. While neuroscience research in developmental cognition and psychopathology is scientifically compelling to many clinicians, few practitioners expect that neuroscience discoveries contribute to and/or modify their day-to-day clinical practice. This science to practice gap is particularly palpable in adolescent mental health, where most established behavioral treatments have less robust outcomes as compared to adults. While increasing levels of attention are being paid to integrative approaches to cross the clinical-neuroimaging divide [2-4], particularly with adolescents [5-7], few concrete, working collaboratives have been advanced yet. Thus, at this time, more publications, research presentations, and communication around this topic must occur in order for this conversation to effectively move forward into the arena of active and productive cooperation and dissemination. This commentary offers one step in this direction by articulating the relevance of this message, which represents a priority area at the National Institutes of Health, and by offering concrete steps to begin engaging in this type of clinical – neuroimaging partnership. Ultimately, this commentary offers suggestions to enhance cross-discipline collaboration, with the goal of improving adolescent treatment outcome.

Adolescence as a time of neural responsiveness and risk

Research is steadily unraveling the relation between the developing brain and emerging psychopathology in adolescence. Described as a “sensitive period,” [8] this time of catalytic neuronal change contributes to increased opportunity and risk. In terms of benefit, the adolescent brain has an elevated capacity to encode and synthesize new information [9]. For

example, experimentation with new behaviors (e.g., driving) combines with plasticity to yield more efficient brain function (e.g., synaptic pruning affected by experience). In this respect, the adolescent brain is relatively more adaptive than during other developmental periods, as it actively receives and synthesizes novel information to improve individual functioning.

Yet, the same evolutionary drive that steers adolescents towards experimentation also enhances the potential for risk. Supporting their predisposition toward reward-seeking, the activated dopaminergic (DA) pathways of the brain encompass regions including the orbitofrontal cortex (OFC), medial prefrontal cortex (mPFC), nucleus accumbens (NAc) and ventral striatum (VS). Largely modulated by the ventral tegmental area (VTA), which has DA projections to these and other salient regions (e.g., amygdala), these networks underlie adolescents' estimation of rewards, along with salient social information. It is argued that the greater activation of these regions during adolescence reflects the redistribution of DA receptor density in these regions that occurs during these years. Relative to other age periods, adolescents subsequently experience a greater release of DA in response to rewarding events. Practically, this means that risk behaviors, often inherently exciting, frightening, and fun, may in fact feel much more so during adolescence. Importantly, many of these same developing mesocorticolimbic systems are also key substrates in the later manifestation of psychopathology, including addiction, during adulthood [10]. Related, the animal and human literatures suggest that one potential modulator of psychopathology onset and course is exposure to common, neurotoxic substances (including alcohol, tobacco, cannabis) during adolescence.

The gap between neurodevelopment and direct clinical practice

Despite strong empirical support for these neurobiological models, many clinicians, including those in addiction, report that these discoveries feel disconnected from their day-to-day practice. At the same time, advances in adolescent psychotherapy research have slowed

[11]. Consequently, clinical researchers are largely at a loss to explain modest expectations for long-term improvement from treatment [e.g., less robust effect sizes for adolescents (ages 12-23; $d = 0.17$) [12] as compared with adults (ages 16-62; $d = 0.77$) [13] 12 months following treatment]. Leading treatment researchers have called for change, “moderate effectiveness calls for improvement and scarce resources call for efficiency...more nuanced analytic methods are...needed” (p. 883) [14].

Despite the commonly-held belief that the brain is at the source of human change, few clinical research teams have looked to the adolescent brain to identify new treatment targets or metrics of outcomes. Understanding how and why the adolescent brain does (or does not) change in the context of treatment might lead to improvements in current treatment approaches such as promoting positive brain response (e.g., greater neural control; activation of contemplation networks). Adolescent brain data offers one promising route to enhance current evidence-based treatments for this high-need, and often underserved age group.

Bridging adolescent neuroscience and treatment

Cutting-edge brain imaging methodologies are a highly-sensitive set of tools to empirically explore neural substrates underlying successes and failures of current clinical treatments. Beginning with more fundamental association studies of brain structure and function [15], many treatment teams are now evaluating *how* adult and adolescent brains respond to treatment. For example, in the context of addiction, initial explorations with adults have evaluated brain response to pharmacotherapies.

Arguably, these explorations may be even more salient and important to the advancement of improvements in the context of behavioral treatment. Neuroimaging data are critical in clinical research so that clinicians and scientists can fully understand the mechanisms underlying treatment successes and failures. Specifically, at this time, our behavioral metrics of adolescent treatment response (e.g., reward response) are not sufficiently sensitive to guide

clinical decision making. Thus, with brain data in hand, we might learn that a particular behavioral treatment (e.g., contingency management) dampens adolescents' neural reward response to drug cues. This information could directly inform clinical decision making, such as determining whether to enhance this behavioral treatment (e.g., contingency management) with medication and/or to include another adjunctive behavioral treatment that has gained empirical support in dampening adolescent neural reward response. Further, through this approach, one might learn that one element of reward neurocircuitry is more plastic and responsive to behavioral treatment than another. Moreover, this approach might uncover that different treatment elements (e.g., motivationally-focused vs. reward-focused behavioral treatments) have different neural targets. Ultimately, learning how clients' brains do or do not respond to these treatment elements could guide us to the use of one treatment target over another. Finally, querying the response of the adolescent brain to different treatment approaches might uncover which treatments (e.g., behavioral approaches vs. medication vs. their combination) have the most enduring effects, and in which key target regions. Together, structural and functional neuroimaging will generate neural targets that can concretely help clinical researchers strengthen existing treatment options. Understanding the biological mechanisms of behavioral change is fundamental to advance growth and make substantive advances in the field of adolescent addiction treatment.

In terms of the clinical-neuroscience divide, novel examinations have begun to evaluate the neural substrates of in-session clinical exchanges (client change talk; therapist statements) by examining functional brain response [16, 17]. By replaying in-session clinical excerpts back to individuals within the scanner, Feldstein Ewing and colleagues found that human brains respond differently to the clients' own in-session statements in favor of change (e.g., *"I need to cut back on smoking weed"*) when contrasted with their own in-session statements in favor of staying the same (e.g., *"I like smoking weed – it's fun!"*). Notably, the location of functional brain response varied by developmental period: mesocorticolimbic reward regions for adults (ages

35-52) [17], and introspection/contemplation areas for adolescents (ages 14-17) [16]. Further, adolescents who showed greater brain response in the precuneus, cingulate gyrus, posterior cingulate, and globus pallidus showed greater behavior change, as measured by significantly less cannabis use post-treatment [16]. Collectively, these studies illuminate the neural processes behind spontaneous and self-generated client thought, both in-session and after leaving a practitioner's office. Other studies are exploring the role of self-generated thought [18], the relationship between brain response and relapse [19], and neurofeedback [20]. Together, these studies reveal that neuroimaging can unlock a previously unavailable empirical window, allowing investigators to access, observe, and understand critical brain processes underlying within- and post-treatment behavior change.

Neuroimaging in the context of psychotherapy for adolescent psychopathology

Although a handful of functional neuroimaging modalities exist, concerns about radioactive material (e.g., PET) and accurate localization (e.g., EEG, MEG) can limit their application to treatment contexts. Two modalities, magnetic resonance imaging (MRI) and functional MRI (fMRI) [21], are excellent for use with youth, as they are non-invasive and offer high-resolution localization.

Despite the advantages of MRI and fMRI, they also have limitations [for a review, see 22]. As examples, fMRI sessions have a restricted duration, hampering psychosocial treatment sessions *in* the scanner. fMRI requires multiple trials of behavior (i.e., repetitions of the event of interest) to produce observable and reliable effects; a condition of interest needs to occur a number of times. Thus, active collaboration between clinicians and neuroimagers is requisite to synchronize protocols to yield translational projects that have ecological clinical validity, and that yield usable, empirical data regarding treatment processes in the brain.

One more critical consideration is that neuroimaging studies are, by and large, correlational in nature [4] [23]. Much of the neuroimaging literature has hinged on examining

correlations, or comparisons of regions and relationships of interest within or between small samples. Ultimately, to generate maximal benefit to advancing our understanding treatment *mechanisms*, the field must move towards causation, by generating empirical data that identify which neuropsychological systems are necessary to successful treatment response (behavior change) [4]. Given attention to issues of replicability and reproducibility in the field of psychological science at this time [24], many labs are currently working on methods, including setting aside a small set of subjects for validation purposes (e.g., “training” vs. “test” sets), to address and minimize prediction errors that stem from use of small datasets in these evaluations (e.g., issue of “optimism”) [23] [25]. It is also important to note that some have found that neuroimaging data has yielded greater predictive power than behavioral metrics alone [23] [26]. One caveat here is that group differences in neural response could represent either the neural systems that are impacted by the disorder itself (e.g., the characterizing features of the disorder), or the mechanisms by which treatment exerts its response (e.g., the areas of the brain that are impacted by the treatment, and that may subserve improved behavioral response and change) [23]. In some cases, these areas may be regions that are related to the etiology and progression of the disease, rather than treatment response. Moreover, these may also be neural regions that have no intuitive theoretical link to the disorder, its treatment response and/or its resolution. Ultimately, these issues represent the crux of the problem wherein “data has been disconnected from practice”. These conversations and considerations are precisely the types of discussions that neuroscientists must have with clinicians in order to advance the field.

Another common problem is that typical and widely-occurring individual and environmental differences can interfere with interpretation of scan data (e.g., left-handedness; past-hour tobacco use). These variables may extend from the most fundamental of variables (e.g., time of day, degree of sleepiness; whether or not a child has eaten, brought their corrective lenses, consumed caffeine, refined sugar, and/or exercised prior to the scan session)

to more intransigent variables that could impact fMRI and structural data, including pre-existing genetic, neurological and/or developmental differences in brain function and structure (e.g., pubertal development status and menstrual cycle; natural differences in age and growth and related cognitive development). Further, there are inherent risks in using functional, task-dependent metrics, where approaches to measures of attention, for example, can vary widely by lab (e.g., anti-saccade vs. GoNoGo; various versions of GoNoGo); in this respect, structural and resting state measures offer some more technical consistency that can improve generalizability of scientific approach and subsequent empirical findings [23].

Limiting these confounding factors is often considered necessary to isolate the true neural signal within the therapeutic effect of interest. Yet, this can inadvertently homogenize samples and their generalizability to broader clinical and non-clinical populations. While many developmental neuroscience teams exclude youth with co-occurring Axis I disorders (outside the target disorder) and common psychotropic medicines (e.g., antidepressants; anxiolytics), the recommendation here is to *include* these samples, and empirically compare brain response and treatment outcomes [e.g., between youth who have comorbid substance use disorder (SUD) and attention deficit hyperactivity disorder (ADHD) vs. youth who only have SUD]. Collaborating with clinical investigators expert on which disorders or patient populations may be integrated (vs. those that would render the data un-interpretable) is key.

Currently, it is straightforward to evaluate brain change in relation to behavior, for example, relative levels of BOLD signal pre- vs. post-treatment to pictures of spiders in the treatment of specific phobia (M age = 24.15) [27]. However, in some contexts, it might be more important to evaluate a more elusive aspect or mechanism of the behavior disorder (such as the nature of substance use – e.g., craving vs. self-regulation - within adolescent addiction, as evaluated with samples of 14-17 year olds [16]). The cognitive neuroscience literature is replete with sophisticated tasks to measure the clinical target of interest (e.g., social processing; theory of mind; attention; distraction; behavioral activation; self-thought).

Further, methodological approaches to evaluate adolescent brain volume and integrity of component tracts evaluate the brain at rest, providing critical information regarding the “wiring” of the adolescent brain even when youth are not behaviorally or cognitively engaged. Disaggregation of treatment components to identify active ingredients linked to these types of structural change may provide additional clues about more enduring brain responses to treatment. For example, youth who receive a coping skills training module might show strengthened white matter connectivity between the prefrontal cortex and subcortical areas as compared to youth who receive no treatment. Research that examines variation across extent of treatment exposure (e.g., number of sessions) might identify the crucial duration necessary for improved connectivity (and therefore possibility of lasting treatment response and decreased rates of high-risk behavior). Recent studies have also been examining how mindfulness may enhance emotion regulation or dampen affective in-the-moment decision making (ages 18-24) [28]; through a synergistic examination, scientists could determine whether this treatment is causally linked to more developmentally-mature and/or “healthier” adolescent brain structure. If so, additional exercises that improve brain health may maximize the benefits of this treatment.

Practical recommendations regarding how data from neuroimaging studies can enhance current adolescent evidence-based treatments.

Improved clinical outcomes informed by neuroscience will ultimately depend on an iterative and integrated series of highly controlled protocols with studies of neural substrates of clinically relevant specific behaviors (e.g., braking when driving fast among adolescents rated consistently as trait-impulsive; ages 14-18) [29]. Researchers may be well-advised to identify their clinical constructs of interest, disaggregate them behaviorally and neurally, and after identifying structural and functional correlates of isolated contributing behaviors, test whether the combination relates to the more clinically relevant target. To accomplish this, collaboration between clinicians and neuroscientists will be paramount.

In **Figure 1**, five steps are recommended, with data from Dr. Feldstein Ewing's ongoing line of empirical research as a tangible example of how these efforts could be applied.

Step 1 is in line with recent recommendations by Allen and Dahl [30], wherein the first foundational step in this type of collaborative is to identify the “modifiable” elements of treatment response that are of interest to the investigative team. Here, the research team must select which factors they believe may be fundamental to an adolescent's positive treatment response (or lack thereof). For example, both behavioral and neuroimaging work by Feldstein Ewing and colleagues has found that certain in-session therapist behaviors positively predict youth treatment response (ages 13-19) [31, 32]. Thus, for this step, one proposed modifiable target could be “in-session therapist behaviors.” It is suggested that this treatment element (therapist behaviors) may be able to directly enhance adolescents' process of self-examination and potentially self-regulation. Importantly, this is a treatment target that is highly amenable and responsive to modification (e.g., through clinical supervision and didactics).

Step 2 revolves around identifying what the research team believes are the developmentally-relevant mechanisms of positive treatment response, both in terms of neurobiology and behavior. Here, Feldstein Ewing and colleagues have found that for substance-using youth, activation of brain networks involved in introspection and contemplation predicted adolescents' successful treatment response (ages 14-19) [16, 32]. Therefore, for this model, one might expect an increase in self-reflection across both neural metrics of self-reflection (e.g., greater network activation), as well as in behavioral estimates of self-reflection (e.g., self-report ratings of self-reflection and contemplation).

Step 3 includes identifying a concrete treatment outcome that is sensitive and appropriate for this developmental period. In adolescent addiction, the recommendation is to move away from a focus on relapse as the metric of successful treatment response, which is widely the outcome of interest in adult studies (e.g., percent days abstinent). Because substance use is normative and typical in this age range [33], abstinence is rarely a clinically

useful (or tenable) treatment target. Instead, when treating adolescents, many clinicians target consumption at less hazardous levels and in less harmful ways (e.g., harm reduction) [34].

Step 4 includes rigorously subjecting these theories to empirical examination by developing neuroimaging and behavioral paradigms that directly test the association of these proposed mechanisms with youth psychosocial treatment response [4]. While not always feasible, the strongest design approaches [4] are ones that include randomization to active control treatment conditions (e.g., motivational interviewing vs. cognitive behavioral therapy), and active comparisons within the neuroimaging paradigm (e.g., therapist complex reflections vs. therapist closed questions) [32]. Pre- post-designs, which examine the modifiable factor prior to treatment exposure and following treatment outcome, are critical to obtain a temporally accurate estimate of how treatment affected the proposed neural mechanism of change [35]. Once these data have been generated, updating the research team's theoretical model to reflect treatment outcomes and the model of treatment response is critical. Feldstein Ewing and colleagues actively empirically test and use these data to refine their theoretical model of treatment response in this very manner [36] [37].

Step 5 is potentially the most important step, which includes leaving the lab to share the translational data with the larger field of clinicians and scientists, to disseminate these findings, and generate dialogue in this area. Feldstein Ewing and colleagues have taken on this challenge by actively publishing their research findings in both clinical as well as neuroimaging journals, discussing empirical outcomes in integrative clinical-neuroscience forums (including www.scienceofchange.org, which targets both clinicians and neuroscientists), sharing the translation of "how these data matter for clinicians" in national and international clinical trainings, and integrating their empirical brain data into their clinical approach and clinical supervision.

In terms of developmental caveats, it is critical to note that the adolescent brain is changing in structure and function throughout this period. Thus, these examinations must occur with an eye to age, gender, and pubertal development status, in order to assess which neural

mechanisms of treatment response are most important, and/or have the most impact at different points in adolescents' developmental trajectory.

Conclusions

There are compelling scientific reasons to merge basic neuroscience with treatment development; however, innovative integrative work can be daunting and difficult. We believe that these practical steps might help form the groundwork for successful collaboration. For example, we encourage teams to begin by engaging in multi-disciplinary collaboration to translate methodologies, data, and interpretations between neuroscientists and clinical researchers. Next, when collaborating across the fields of clinical treatment and neuroimaging, it is integral to learn and translate one another's jargon. By this, we encourage teams to sit down together to ensure that any familiar and unfamiliar terminology has the correct interpretation for both fields. For example, terms that are widely used but mean different things in the neuroimaging and clinical literatures include measures of "behavior" or "behavioral response", and/or "impulsivity". Next, it is encouraged that collaborative teams ask questions, as no well-worn path exists in synergistic clinical neuroscience inquiry. Openly querying any terms that are new or unfamiliar is a worthwhile endeavor, and will help establish successful clinical research collaboration. In this sense, it is important to be comfortable with sounding like a novice in the unfamiliar field. For example, it is okay to query what a "regressor", "covariate" or "controlling for" means in the context of neuroimaging analyses. Fundamentals that are clear to neuroimagers may not be clear to clinicians (e.g., "BOLD response"; "white matter"; "gray matter"; "DTI"; "prepotent response"). Similarly, neuroimagers may not know terms or concrete examples of commonly accepted jargon in clinical communities ("therapist behaviors", "in-session response", "clinically significant change"). Additionally, we must brainstorm methods to effectively pave an open road for communication of study findings between clinicians and neuroimagers. One clear arena to facilitate dynamic communication is via scientific meeting; in

one active effort, the Drs. T Chung and SW Feldstein Ewing convened an annual forum where both fields shared scientific advances (www.scienceofchange.org). This has proven to be a highly fruitful avenue to begin active dialogue and communication, with attention to the practical relevance of findings across the aisle to both fields (clinician to neuroimager; neuroimager to clinician). However, more work is needed to develop an ongoing and accessible platform for neuroimagers to disseminate their findings to clinicians and for clinicians to share their clinical observations to neuroimagers. With this active communication, this type of scientific synthesis will catalyze the field.

Ultimately, understanding brain structure and function subserving treatment response, and using methodologies that more effectively fuse meaningful clinical targets with neuroimaging methodology, should push treatment innovation further. Through these efforts, improved treatment response, both acute and long-term, should result.

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Figure 1**Overview of steps:**

Step 1: Identify developmentally-relevant modifiable treatment target

Step 2: Identify developmentally-relevant treatment mechanism (biology and behavior)

Step 3: Identify developmentally-relevant treatment outcome

Step 4: Empirically evaluate the model. With resultant data, update empirical model of treatment response.

Step 5: Disseminate and train clinicians in updated treatment approach.

Example model: