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*European Journal of Neuroscience*, Vol. 52, No. 11 (December 2020): 4546-4562. [DOI](http://doi.org/10.1111/ejn.14927). This article is © Wiley and permission has been granted for this version to appear in [e-Publications@Marquette](http://epublications.marquette.edu/). Wiley does not grant permission for this article to be further copied/distributed or hosted elsewhere without express permission from Wiley.

Organic Cation Transporter 3 And the Dopamine Transporter Differentially Regulate Catecholamine Uptake in The Basolateral Amygdala and Nucleus Accumbens.

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# Abstract

Regional alterations in kinetics of catecholamine uptake are due in part to variations in clearance mechanisms. The rate of clearance is a critical determinant of the strength of catecholamine signaling. Catecholamine transmission in the nucleus accumbens core (NAcc) and basolateral amygdala (BLA) is of particular interest due to involvement of these regions in cognition and motivation. Previous work has shown that catecholamine clearance in the NAcc is largely mediated by the dopamine transporter (DAT), but clearance in the BLA is less DAT‐dependent. A growing body of literature suggests that organic cation transporter 3 (OCT3) also contributes to catecholamine clearance in both regions. Consistent with different clearance mechanisms between regions, catecholamine clearance is more rapid in the NAcc than in the BLA, though mechanisms underlying this have not been resolved. We compared the expression of DAT and OCT3 and their contributions to catecholamine clearance in the NAcc and BLA. We found DAT protein levels were ~ 4‐fold higher in the NAcc than in the BLA, while OCT3 protein expression was similar between the two regions. Immunofluorescent labeling of the two transporters in brain sections confirmed these findings. Ex vivo voltammetry demonstrated that the magnitude of catecholamine release was greater, and the clearance rate was faster in the NAcc than in the BLA. Additionally, catecholamine clearance in the BLA was more sensitive to the OCT3 inhibitor corticosterone, while clearance in the NAcc was more cocaine sensitive. These distinctions in catecholamine clearance may underlie differential effects of catecholamines on behavioral outputs mediated by these regions.

# Keywords:

cocaine; corticosterone; dopamine; release; uptake; voltammetry

OCT3 has a greater role in catecholamine uptake in the BLA than in the NAcc. The DAT is more highly expressed in the NAcc than the BLA, whereas OCT3 is expressed similarly between the regions. In the BLA, uptake is far more inhibited by corticosterone (an OCT3 blocker) than it is within the NAcc.Abs.

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# Abbreviations

* ACSF artificial cerebrospinal fluid
* BLA basolateral amygdala
* CA catecholamine
* CORT corticosterone
* DAT KO dopamine transporter knockout mice
* DAT dopamine transporter
* NAcc nucleus accumbens core
* NET norepinephrine transporter
* OCT3 organic cation transporter 3
* PB phosphate buffer
* PBS phosphate‐buffered saline
* PMAT plasma membrane monoamine transporter

# [INTRODUCTION](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

The nucleus accumbens core (NAcc) and basolateral amygdala (BLA) receive dopaminergic projections from the ventral tegmental area (VTA) that have been implicated in regulating aspects of cognition (Churchwell, Morris, Heurtelou, & Kesner, 2009; Fuxe, Borroto‐Escuela, Romero‐Fernandez, Zhang, & Agnati, 2013; Li, Zuo, Yu, Ping, & Cui, 2015; Touzani, Bodnar, & Sclafani, 2008; Perez de la Mora, Jacobsen, Crespo‐Ramirez, Flores‐Gracia, Fuxe, 2008), fear (Klumpers, Morgan, Terburg, Stein, & van Honk, 2015; Li & McNally, 2015; Stevenson & Gratton, 2004), and motivated behaviors (Ballester Gonzalez, Dvorkin‐Gheva, Silva, Foster, & Szechtman, 2015; Shiflett & Balleine, 2010; Simmons & Neill, 2009). Though both NAcc and BLA are implicated in these behaviors, dopamine signaling seems to have opposite effects in the two areas. Dopamine neurotransmission in the NAcc has been historically linked to the psychomotor stimulating and rewarding effects of drugs of abuse (Baik, 2013; Fish, DiBerto, Krouse, Robinson, & Malanga, 2014; Nakamura et al., 2014), and functional reduction of dopamine transmission in this region has been observed following chronic drug exposure and withdrawal (Markou & Koob, 1992; Rose et al., 2016; Schulteis, Markou, Cole, & Koob, 1995; Siciliano, Calipari, Ferris, & Jones, 2015; Siciliano et al., 2016). The behavioral effects of dopamine in the BLA are often the opposite. For example, augmented dopamine in the NAcc is linked to positive‐affective states (Baik, 2013; Fish et al., 2014; Nakamura et al., 2014), while increased dopamine in the BLA is associated with negative affect or anxiogenic states (Bananej, Karimi‐Sori, Zarrindast, & Ahmadi, 2012; Diaz, Chappell, Christian, Anderson, & McCool, 2011; Karkhanis, Rose, Huggins, Konstantopoulos, & Jones, 2015). Further, dopamine is decreased in the NAcc following chronic adolescent social isolation in rats, a model of early life stress, whereas dopamine release is increased in the BLA using the same paradigm (Karkhanis et al., 2015).

Catecholamine neurotransmission can be influenced by both release and clearance mechanisms, which can differ widely between brain regions. Multiple mechanisms of dopamine clearance have been delineated, including uptake by both high‐affinity (Budygin, John, Mateo, & Jones, 2002; Gainetdinov, Jones, & Caron, 1999; Gainetdinov, Jones, Fumagalli, Wightman, & Caron, 1998) and low‐affinity transporters (Baganz, Horton, Martin, Holmes, & Daws, 2010; Duan & Wang, 2010; Gasser, Lowry, & Orchinik, 2006; Hill & Gasser, 2013; Lips et al., 2005), in addition to enzymatic degradation (Akil et al., 2003; Napolitano, Bellini, Borroni, Zurcher, & Bonuccelli, 2003) and diffusion (Kehr, Hoistad, & Fuxe, 2000; Trouillon, Lin, Mellander, Keighron, & Ewing, 2013). Dopamine clearance is regulated by distinct mechanisms in the NAcc and BLA. Dopamine transporters (DAT), are high‐affinity, low‐capacity transport proteins abundantly expressed in the NAcc (Karkhanis et al., (2015); Gainetdinov et al., 1998; Gainetdinov et al., 1999; Budygin et al., 2002), but are less abundant in the BLA. A growing body of literature suggests that the organic cation transporter 3 (OCT3) (Baganz et al., 2010; Duan & Wang, 2010; Gasser, 2018; Gasser, Hurley, Chan, & Pickel, 2017; Gasser et al., 2006; Graf et al., 2013; Hill & Gasser, 2013; Iversen & Salt, 1970; Lightman & Iversen, 1969; Wieland, Hayer‐Zillgen, Bonisch, & Bruss, 2000), which is expressed in both the NAcc (Graf et al., 2013) and BLA (Hill & Gasser, 2013), can mediate uptake of a variety of monoamine species including serotonin (Baganz et al., 2010; Schmitt et al., 2003), norepinephrine (Duan & Wang, 2010), and dopamine (Gasser et al., 2006; Hill & Gasser, 2013). Compared to the DAT, OCT3 has a lower affinity for dopamine (Duan & Wang, 2010), but a higher capacity for dopamine transport (Zhu, Appel, Grundemann, & Markowitz, 2010). The relative contributions of OCT3 and the DAT to dopamine clearance in the NAcc and BLA have not been directly investigated. Elucidating this relationship could provide unique pharmacological treatment strategies for ameliorating substance use disorders and anxiety‐related disorders by regulating distinct dopamine transmission in the NAcc and BLA.

Exposure to stressors engages the hypothalamic–pituitary–adrenal (HPA) axis (Franco et al., 2016; Hek et al., 2013) to increase synthesis and release of corticosteroid hormones (predominantly cortisol in humans, and corticosterone (CORT) in rodents) (Jones, Gupton, & Curtis, 2016; Jung et al., 2014; Ottenweller, Natelson, Pitman, & Drastal, 1989; Smyth et al., 2015). In the NAcc, CORT decreases the rate of dopamine clearance (Graf et al., 2013) and thereby increases the duration of neurotransmitter action in this region. CORT and other corticosteroids acutely and directly inhibit OCT3‐mediated transport (Baganz et al., 2010; Wieland et al., 2000); in fact, recent work has demonstrated that the pharmacological interaction between CORT and OCT3 (Graf et al., 2013) is functionally analogous to uptake inhibition by cocaine of the DAT (Grundemann, Schechinger, Rappold, & Schomig, 1998). CORT treatment, likely by inhibiting OCT3‐mediated clearance, has been shown to augment extracellular concentrations of serotonin (Baganz et al., 2010), norepinephrine (Ayala‐Lopez et al., 2015) and dopamine (Graf et al., 2013). Taken together, this suggests that OCT3 blockade may augment extracellular catecholamine levels (Baganz et al., 2010; Duan & Wang, 2010; Gasser et al., 2006; Hill & Gasser, 2013; Schmitt et al., 2003). Notably, OCT3 is densely expressed in the BLA (Hill & Gasser, 2013) where CORT may exert similar inhibitory effects upon OCT3 to decrease catecholamine clearance. By regulating the duration of released catecholamines, clearance mechanisms contribute greatly to the extent to which receptors are activated on target cells and thus can have powerful effects on behavior modulated by catecholamines.

Previous work has shown that the catecholamine clearance rate in the NAcc is significantly faster than clearance rates in the BLA (Jones, Garris, Kilts, & Wightman, 1995). It is plausible that this distinction is due to differences in both the specific transporters expressed, and the density of transporter expression in the two areas. Our laboratory has previously shown that DAT protein expression is lower in the BLA than in the NAcc (Karkhanis et al., 2015) and that the apparent affinity of transporters for catecholamines is lower in the BLA compared to the NAcc (Jones et al., 1995). Together, these factors likely contribute to regional differences in clearance rates. In consideration of these data, we examined the localization and expression levels of DAT and OCT3 in the NAcc and BLA using immunofluorescence and Western blot and assessed the functional contributions of DAT and OCT3 to catecholamine clearance in the two regions of drug‐naïve rats using ex vivo fast‐scan cyclic voltammetry.

# [METHODS](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

## [Experimental animals](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

Male Sprague‐Dawley rats (*n* = 6–10 per group, 9–10 months old, Harlan Laboratories, Indianapolis, IN) and homozygote DAT knockout (DAT KO) mice were used for experiments. DAT KO mice were bred in‐house via crossing heterozygous DAT+/− 129SvJ and C57BL/6 mice (Lack, Jones, & Roberts, 2008), and then breeding resultant DAT+/− mice. Tissue samples were obtained from DAT+/− × DAT+/− breeding pair litters and digested prior to phenol:chloroform DNA extraction. Polymerase chain reaction was run with forward and reverse oligonucleotide primers for DAT to verify animals with full DAT KO. All animals were maintained on a 12:12 light cycle and given standard rodent chow and water ad libitum, and all experiments were performed in the dark phase of the animals' light cycle. Animal care and handling, as well as experimental protocols, were approved by the Wake Forest School of Medicine Institutional Animal Care and Use Committee and adhered to all National Institutes of Health Animal Care Guidelines.

## [Brain slice preparation for fast‐scan cyclic voltammetry](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

Animals were deeply anesthetized with isoflurane gas in an induction chamber and then rapidly decapitated. Brains were removed and placed into ice‐cold, pre‐oxygenated (95% O2/5% CO2) artificial cerebrospinal fluid (ACSF; in mM: 126 NaCl, 2.5 KCl, 1.2 NaH2PO4, 1.4 CaCl2, 2.4 MgCl2, 25 NaHCO3, 11.0 glucose, 0.4 l‐ascorbic acid; pH adjusted to 7.4). A vibrating tissue slicer (Leica Biosystems, Buffalo Grove, IL, USA) was used to prepare coronal brain slices (400µm thick from rats, 300 µm thick from mice) containing the NAcc (both rats and mice) and BLA (rats only). NAcc and BLA tissue for protein analysis was excised from slices by hand.

## [Western blot hybridization](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

DAT and OCT3 protein levels were quantified using Western blot hybridization following methods described previously (Giros, Jaber, Jones, Wightman, & Caron, 1996), with slight alterations for OCT3 blots described below. Tissue samples from rats were homogenized in radioimmunoprecipitation assay (RIPA) buffer (150 mM NaCl, 1.0% Triton X‐100, 0.5% sodium deoxycholate, 0.1% sodium dodecyl sulfate (SDS), 50 mM Trizma Base, pH 8.0) and centrifuged at 12,000 *g* for 30 min. Protein concentrations were determined using a BCA protein assay kit (Thermo Scientific, Rockford, IL) and Molecular Devices Spectra Max 384 Plus spectrophotometer (Sunnyvale, CA) running SoftMax Pro software. A MagicMark™ XP molecular weight ladder (LC5602; Thermo Fisher Scientific, Grand Island, NY) and 30μg of protein were loaded into 4%–12% NuPAGE Bis‐Tris gels (NP0321BOX; Thermo Fisher Scientific). Proteins from the gel were transferred onto a polyvinylidene difluoride membrane and probed using the following primary antibody dilutions: DAT (1:4,000; 2,231, EMD Millipore, Billerica, MA), OCT3 (1:1,000; (OCT31‐A, Alpha Diagnostic Int., San Antonio, TX), and β‐actin (1:4,000; ab8229, Abcam, Cambridge, MA). A horseradish peroxidase (HRP)‐conjugated secondary antibody was used (1:5,000 for DAT and β‐actin, 1:4,000 for OCT3) (65‐6120, Thermo Fisher Scientific, Grand Island, NY), in combination with Pierce ECL chemiluminescence (32,106, Thermo Fisher Scientific, Grand Island, NY). Blots were exposed using a Bio‐Rad ChemiDoc imaging system, and the band signal intensity was assessed using QuantityOne software (Bio‐Rad, Hercules, CA).

## [Perfusion and histology](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

Rats were deeply anesthetized by intraperitoneal injection of sodium pentobarbital (100 mg/kg) and were transcardially perfused with ice‐cold 0.05 M phosphate‐buffered saline followed by 4% paraformaldehyde in 0.1 M sodium phosphate buffer (PB, pH 7.4). Following perfusion, brains were removed and post‐fixed in the 4% paraformaldehyde solution for 12 hr at 4°C and were rinsed twice in 0.1 M PB for 12 hr. The brains were then incubated in 30% sucrose in 0.1 MPB for approximately 72 hr. Brains were then blocked into two pieces with a cut in the coronal plane at the caudal border of the mammillary bodies (approximately –5.30 mm bregma) using a rat brain matrix (RBM‐4000C, ASI Instruments, Warren, MI, USA). Brains were frozen rapidly in dry‐ice‐chilled liquid isopentane and stored at −80°C until sectioning. Forebrain sections (25 µm), including the BLA, were cut across the coronal plane using a cryostat (Leica Biosystems, Buffalo Grove, IL, USA) and stored as six alternate sets of sections in cryoprotectant (30% ethylene glycol (w/w)/20% glycerol (w/w) in 0.05 MPB, pH 7.4) at −20°C until immunostaining.

## [Antibodies](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

For immunodetection of OCT3, an affinity‐isolated antibody (RRID AB\_1622571, rabbit anti‐OCT3, cat # OCT31A, Alpha Diagnostics International, San Antonio, TX, USA) raised against an 18‐amino acid sequence in the large intracellular loop of rat OCT3 (amino acids 313–330: HLSSNYSEITVTDEEVSN) was used. This amino acid sequence is 100% conserved between mouse and rat OCT3 and has no significant sequence homology with other organic cation/carnitine transporters. The specificity of this antibody was confirmed previously in immunohistochemical and immunofluorescence applications (Gasser et al., 2006; Gasser, Orchinik, Raju, & Lowry, 2009; Lips et al., 2005; Yorgason et al., 2016). For immunodetection of DAT, an affinity‐purified antibody (RRID AB\_1586991, rabbit anti‐DAT, cat # AB2231, MilliporeSigma, Burlington, MA, USA) was used at a dilution of 1:500.

## [Immunofluorescence](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

Separate coronal sections (25 µm thick) containing BLA and NAcc were used for detection of OCT3 and DAT. After rinsing in PBS, sections were incubated overnight with anti‐OCT3 antibody (1:400) or anti‐DAT antibody (1:500) in phosphate‐buffered saline with 0.1%Tween 20 (PBST). Sections were rinsed the next day and incubated for 2 hr with fluorophore‐conjugated secondary antibodies (AlexaFluor594‐conjugated donkey anti‐rabbit; 1:2000; Jackson ImmunoResearch, West Grove, PA, USA). Sections were then rinsed briefly in PB, mounted onto SuperFrost microscope slides, dried, and coverslipped with EverBrite antifade mounting medium (Biotium, Fremont, CA, USA).

## [Imaging](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

Photomicrographs were acquired using a Nikon 80i microscope fitted with an ORCA‐Flash 4.0LT digital camera (Hamamatsu, Japan) linked to a computer running NIS Elements‐BR software (Nikon Instruments, Melville, NY).

## [Ex vivo fast‐scan cyclic voltammetry](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

Fast‐scan cyclic voltammetry was used to characterize catecholamine terminal function in the NAcc and BLA. Brain slices were transferred to testing chambers and incubated in oxygenated ACSF, heated to 32°C for one hour prior to experiment start. A glass capillary containing a carbon fiber (house‐made; ≈100 µM length, 7 µM radius; Goodfellow Corporation, Berwyn, PA) and a bipolar stimulating electrode (Plastics One, Roanoke, VA) were placed in close proximity (≈100 µm) on the surface of the slice. Endogenous catecholamine efflux was induced by a single‐pulse (350 µA, 4 msec, NAcc) or 10‐pulse stimulation (20Hz, 350 µA, 4 msec/pulse, BLA), applied every five or seven minutes, respectively. Extracellular catecholamine concentrations were detected by applying a triangular waveform (−0.4 to +1.2 to −0.4 V vs. silver/silver chloride, 400 V/sec) every 100 milliseconds to the recording electrode and measuring changes in current at the oxidation potential of dopamine and norepinephrine (~0.6 V). Pharmacological agents (cocaine and/or CORT; as described in Results) were applied once a stable baseline was achieved (three consecutive stimulations within 10% of the previous collection). Electrically evoked catecholamine concentrations were assessed by comparing the current at the peak oxidation potential for dopamine and norepinephrine to each electrode's calibration to known concentrations of catecholamine (3.0μM).

Baseline catecholamine release ([CA] per pulse, [CA]/pulse) and the maximal rate of uptake at catecholamine transporters () were analyzed to assess release and uptake (Wightman *et al*., 1988). Baseline release and uptake were assessed in every slice that was used for recordings, typically 2–4 slices per animal. Standard Michaelis–Menten modeling procedures were used to assess kinetic parameters of baseline stimulation before application of cocaine or CORT. [CA]/pulse was calculated as the amount of catecholamine released per electrical stimulation, and catecholamine uptake was determined with and T80, the time it takes for [CA] to decay to 80% of its peak. *K*m was set to 160 nM for each slice in both regions, based on the known affinity of dopamine for DATs (Vialou, Amphoux, Zwart, Giros, & Gautron, 2004) while pre‐drug values were allowed to vary. Because the affinity of transporters for their endogenous monoamine neurotransmitter (*K*m) varies across transporter species, the use of apparent *K*m (the apparent alteration in affinity of a transporter for its neurotransmitter based on blockade of that transporter, e.g., inhibition of DAT function with cocaine application) was not possible in these experiments, given the examination of multiple transporter species. Therefore, uptake in the NAcc and BLA was assessed using T80, the time it takes for a catecholamine trace to return to 80% of its peak height, a measure of clearance rate. As dopamine and norepinephrine are released in the BLA, "catecholamines" (CA) will be used to describe neurotransmission within this region. All data were collected and analyzed with Demon Voltammetry and Analysis software (Yorgason, Espana, & Jones, 2011).

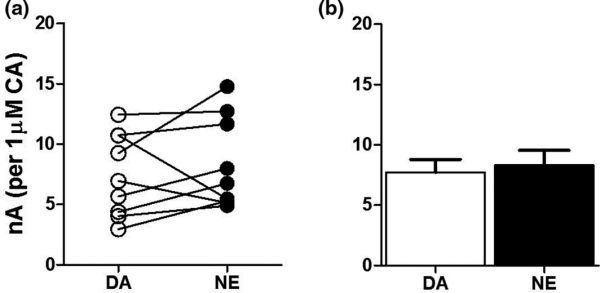
## [Statistics](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

GraphPad Prism (version 7, La Jolla, CA, USA) was used for all statistical analyses and to prepare all graphs. Student's *t* tests were used to determine regional differences in DAT and OCT3 protein levels, baseline [CA]/pulse and clearance measures, effects of maximal concentrations of cocaine or CORT, and the relative contribution of the DAT and OCT3 in uptake. A two‐way repeated measures analysis of variance (ANOVA), with region and drug concentration as factors, was used to determine differences in cocaine and CORT potency between regions. In the event of significant main effects, Bonferroni *post hoc* analysis was used to determine significant group differences. A one‐way ANOVA was used to calculate the effects of CORT on dopamine release in the NAcc of DAT KO mice. Tukey's *post hoc* test was applied when a significant finding was revealed. In voltammetry experiments, 2–4 slices from each animal were used to determine basal catecholamine kinetics (release and uptake; *N* = 15–22 slices/group), whereas one slice per animal was used for drug applications (*N* = 5–10 animals/group).

# [RESULTS](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

## [Carbon fiber microelectrodes have equal sensitivity to dopamine and norepinephrine](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

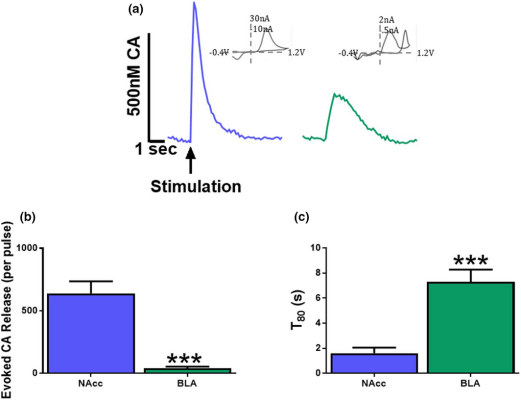
Proper identification of electroactive neurotransmitters using voltammetry relies on neurotransmitter species‐specific cyclic voltammograms, which trace the peak oxidation and reduction potentials of the neurotransmitter. Notably, the oxidation and reduction potentials of dopamine and norepinephrine are nearly identical (Heien, Phillips, Stuber, Seipel, & Wightman, 2003), preventing the separation of these neurotransmitters using voltammetry without the assistance of additional pharmacology. Electrical stimulation of the NAcc causes dopamine efflux (John & Jones, 2007), while stimulation of the BLA causes release of both dopamine and norepinephrine (Brinley‐Reed & McDonald, 1999; Zhang, Muller, & McDonald, 2013). Unfortunately, any pharmacological measures to isolate either dopamine or norepinephrine in the BLA would further reduce the low catecholamine levels measured in this region (Jones et al., 1995) and compromise accurate quantification of stimulated neurotransmitter release. Furthermore, exogenous application of neurotransmitter over the slice does not replicate electrically stimulated endogenous release with reasonable temporal fidelity, and therefore, physiological uptake kinetics would be difficult to ascertain utilizing this method. We sought to discern the sensitivity of our electrodes to each catecholamine to identify any bias to either neurotransmitter that may influence the interpretation of our experiments. To this end, we flowed a moderately high concentration (3 μM) of either dopamine or norepinephrine over the carbon fiber microelectrodes. We chose this concentration in order to more precisely assess sensitivity to the catecholamines through the production of a somewhat larger signal, while by no means reaching the maximum limit of detection. Our data show no difference in electrode sensitivity with respect to dopamine versus norepinephrine (*t*17 = 0.3516, *p* = .729; Figure 1a,b), indicating that the carbon fiber microelectrodes used in the present voltammetry experiments do not favor one catecholamine over the other at the concentration tested.



1 Electrodes have equal sensitivity to dopamine and norepinephrine. (a) Peak oxidation potentials for dopamine (DA) and norepinephrine (NE) showed little variation when detected by the same carbon fiber microelectrode. (b) Within‐electrode analysis revealed no group difference in electrode sensitivity with respect to 3 μM dopamine and norepinephrine . /group

## [Catecholamine release is greater and uptake is faster in the NAcc compared to the BLA](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

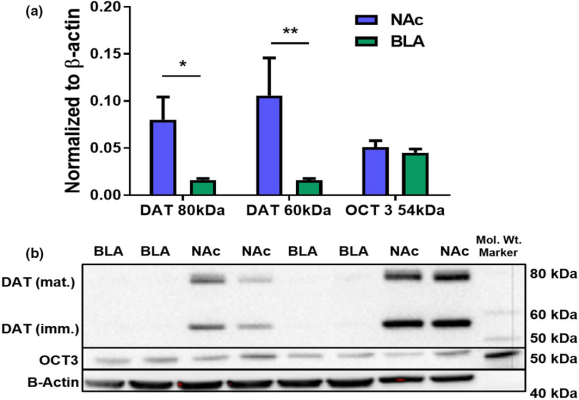
To determine regional differences in catecholamine release and uptake kinetics, pre‐drug catecholamine release and uptake were analyzed. The BLA has reduced catecholamine release compared to the NAcc (Jones et al., 1995; Yetnikoff, Lavezzi, Reichard, & Zahm, 2014), so stronger stimulation parameters were used in the BLA (NAcc, single pulse; BLA, 10 pulse/20 hz). For this reason, release data are analyzed as a measure of catecholamine release per stimulation pulse ([CA]/pulse) rather than overall catecholamine release (µM). Representative traces of these stimulation parameters in the NAcc (blue line) and BLA (green line) are depicted in Figure 2a. Two to four slices from each animal were used to assess release and uptake, prior to subsequent drug application. Student's *t* test revealed significantly reduced [CA]/pulse (*t*19 = 3.558, *p* < .0001, *N* = 22 NAcc, *N* = 17 BLA; Figure 2b) in the BLA compared to the NAcc. Because lower levels of release, particularly within the BLA, may not be great enough to ensure clearance at a saturable level consistently (John et al., 2007), we chose to utilize T80, the amount of time it takes for the voltammetric trace to decay to 80% of its peak height, rather than in order to ascertain reuptake rates. The rate of catecholamine uptake was significantly reduced in the BLA compared to the NAcc as indicated by increased T80 (*t*34 = 5.828, *p* < .0001; *N* = 22 NAcc, *N* = 15 BLA Figure 2c). Additionally, we find that is significantly reduced in the BLA compared to the NAcc (data not shown) demonstrating that the effect on is consistent with T80 data. These data confirm an overall reduction in catecholamine release and uptake dynamics in the BLA.



2 Catecholamine release is greater and uptake is faster in the NAcc compared to the BLA. (a) Representative traces of catecholamine (CA) release and uptake from brain slices containing the NAcc and BLA. (b) Evoked CA release was significantly lower in the BLA compared with to the NAcc (p < .0001). (c) CA uptake, as measured by T80, of the NAcc is significantly faster than BLA slices (p < .0001). NAcc, nucleus accumbens core; BLA, basolateral amygdala; CA, catecholamine. N = 15–22 slices/group from 6–8 animals [Colour figure can be viewed at wileyonlinelibrary.com]

## [DAT protein levels are higher in the NAcc than in the BLA, but OCT3 protein levels are simila](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)r between regions

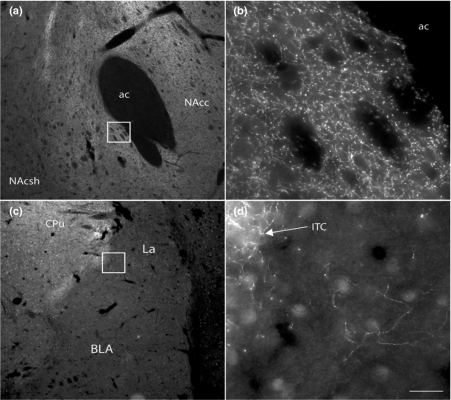
Western blot hybridization was used to compare protein levels of DAT and OCT3 in the NAcc and BLA (*N* = 8 animals/group). DAT protein content, both 80 kDa (functionally mature; *p* = .0247) and 60 kDa (functionally immature; *p* = .0011) isoforms, was significantly higher in NAcc tissue than in BLA tissue (Figure 3a,b). OCT3 protein levels were not different between the two regions (*p* > .05; Figure 3a). Representative Western blots are shown in Figure 3b.



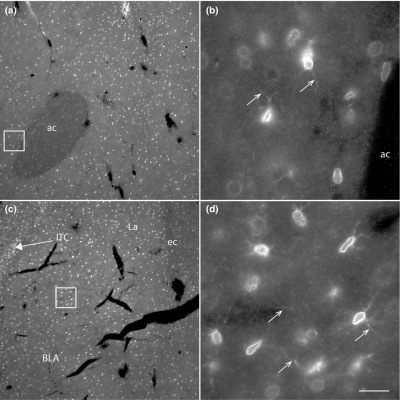
3 DAT protein levels are higher in the NAcc compared to the BLA, but OCT3 levels are similar between regions. Western blot hybridization was used to detect and quantitate the observed variations in DAT and OCT3 densities in the NAcc and BLA. (a) DAT protein levels, normalized to β‐actin, were significantly elevated in the NAcc compared to the BLA at both 78 kDa (p = .0247) and 56 kDa (p = .0011) molecular weights representing the mature and immature DAT isoforms, respectively. Conversely, the protein levels of OCT3 were similar in the NAcc and BLA (p > .05). (b) Representative blot showing NAcc and BLA DAT and OCT3 protein levels quantified in the graph above. NAcc, nucleus accumbens core; BLA, basolateral amygdala; \*p < .05. (N = 8 animals/group) [Colour figure can be viewed at wileyonlinelibrary.com]

## [Localization and density of OCT3 and DAT immunoreactivity in the NAcc and BLA](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

The relative distributions of DAT and OCT3 protein in the NAcc and BLA were examined using immunofluorescence techniques. DAT‐like immunoreactive fibers were observed in both regions; however, they occurred at a much higher density in the NAcc than in the BLA (Figure 4). OCT3‐like immunoreactive perikarya and processes were observed at similar densities in NAcc and BLA (Figure 5).



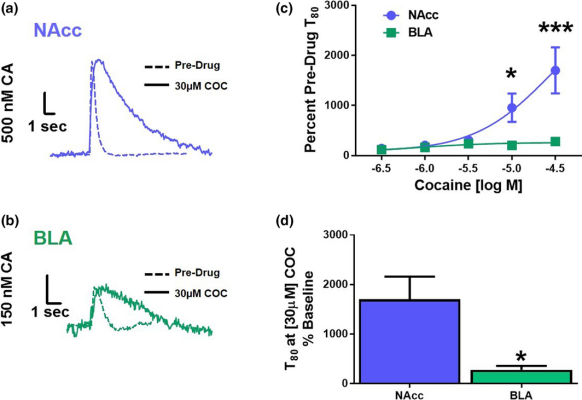
4 Localization of DAT expression in NAcc and BLA. Fluorescence photomicrographs depicting immunofluorescence localization of dopamine transporter (DAT) immunoreactivity in the nucleus accumbens (a, b) and basolateral amygdala (c, d) of the rat. Boxes in (a) and (c) are shown at higher magnification in (b) and (d), respectively. DAT‐immunoreactive fibers were observed at high density in the nucleus accumbens (b) and dorsal striatum (c), and at much lower density in the BLA (d). DAT‐immunoreactive fibers were observed at higher density in the intercalated cell groups (ITC) of the amygdala (c, d) than in the rest of the amygdala. Scale bar = 200 μm (a, c); 20 μm (b, d). ac, anterior commissure; BLA, basolateral amygdala; ITC, intercalated cell group; NAcc, nucleus accumbens core; NAcsh, nucleus accumbens shell



5 Localization of OCT3 expression in NAcc and BLA. Fluorescence photomicrographs depicting immunofluorescence localization of OCT3 immunoreactivity in the nucleus accumbens (a, b) and basolateral amygdala (c, d) of the rat. Boxes in (a) and (c) are shown at higher magnification in (b) and (d), respectively. OCT3‐immunoreactive punctae, perikaryae, and processes (arrows in b, d) were observed at similar densities in both areas. OCT3 perikarya were observed at higher density in the intercalated cell groups (ITC) of the amygdala (c) than in the rest of the amygdala. Arrows in b, d indicate OCT3‐immunoreactive processes. Scale bar = 200 μm (a, c); 20 μm (b, d). ac, anterior commissure; BLA, basolateral amygdala; ITC, intercalated cell group

## [Inhibition of catecholamine uptake by cocaine is greater in the NAcc than the BLA](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

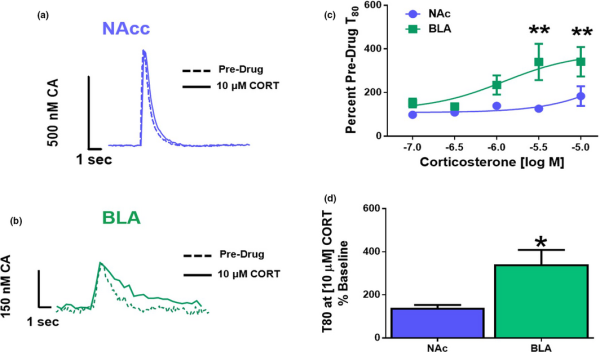
In order to assess regional differences in the DAT‐mediated catecholamine uptake, we examined the effects of cocaine, a DAT inhibitor, on catecholamine clearance in the NAcc and BLA (Figure 6, *N* = 5 per group). Cocaine inhibits catecholamine uptake through DAT and other monoamine transporters (Calligaro & Eldefrawi, 1987; Jones et al., 1995; Uhl, Hall, & Sora, 2002; Yorgason et al., 2011), an effect that is measured as an increase in T80. Figure 6 depicts representative traces collected from NAcc (Figure 6a) and BLA (Figure 6b) slices before and after stimulation under control conditions and following incubation with 30 µM cocaine. Two‐way repeated measures ANOVA comparing the effects of a cumulative cocaine concentration response curve on inhibition of dopamine uptake in the NAcc and BLA revealed a main effect of region (*F*1,33 = 18.90, *p* < .0001; Figure 6c) and drug concentration (*F*4,32 = 8.246, *p* < .0001; Figure 6c) on T80 over the cocaine concentration response curve. Additionally, an interaction between region and drug dose (*F*5,35 = 6.835, *p* < .0008; Figure 6c) was detected. Bonferroni *post hoc* analysis demonstrated a significant difference in uptake inhibition between regions at the 10 µM (*p* < .05) and 30 µM dose (*p* < .001). Comparison of T80 at the highest cocaine concentration as a percentage of pre‐drug T80 in the NAcc and BLA using a Student's *t* test revealed a significantly greater potency of cocaine in the NAcc compared to the BLA (*t*5 = 2.581, *p* < .0494; Figure 6d). Of note is a lack of increased release amplitude in the presence of cocaine, which may seem counterintuitive. However, this is consistent with previous work in our laboratory that demonstrates biphasic effects of dopamine amplitude with increasing concentrations of cocaine. Augmentation of signal amplitude with cocaine administration peaks at approximately the 1–3 µM range; however, we have observed consistently that at greater concentrations, the peak amplitude begins to decline (John & Jones, 2007; Yorgason et al., 2011). At a concentration of 30 µM, cocaine application no longer induces an increase in signal amplitude due primarily to activation of presynaptic autoreceptors (Wieczorek & Kruk, 1994).



6 Inhibition of catecholamine uptake by cocaine is greater in the NAcc compared to the BLA. (a) Representative traces of evoked catecholamine release and uptake in brain slices containing the nucleus accumbens (NAcc) at baseline (Pre‐Drug) and 30 µM cocaine (COC), overlaid. (b) Represented traces from basolateral amygdala (BLA)‐containing brain slices at baseline and 30µM COC overlaid. (c) Cocaine (10 and 30 μM) significantly impaired catecholamine clearance in the NAcc compared to the BLA, (p < .05, and p < .001, respectively). (d) An increase in NAcc T80 revealed a significantly greater potency of COC in the NAcc, compared to the BLA (p < .05). NAcc, nucleus accumbens core; BLA, basolateral amygdala; CA, catecholamine; COC, cocaine; \*p < .05; \*\*\*p < .001. N = 5 animals/group [Colour figure can be viewed at wileyonlinelibrary.com]

## [Inhibition of catecholamine uptake by corticosterone is greater in the BLA than the NAcc](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

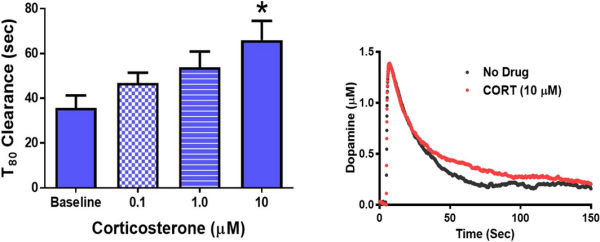
To determine the relative contribution of OCT3 to catecholamine clearance in the NAcc and BLA, we examined the effects of CORT, which inhibits OCT3‐mediated transport, on catecholamine uptake in the two regions. Figure 7 depicts representative traces collected from the NAcc (Figure 7a) and BLA (Figure 7b) before and after stimulation, under control conditions and following treatment with 10 µM CORT. Two‐way repeated measures ANOVA revealed a dose‐dependent increase in catecholamine clearance time, T80, in the BLA across the concentration response curve (*F*4,42 = 3.929, *p* < .0085; Figure 7c). Bonferroni *post hoc* analysis revealed a significant difference in T80 between regions at the 3.0 µM (*p* < .01) and 10 µM (*p* < .01) concentrations. Student's *t* test of group data at the highest CORT concentration tested (10 µM) further underscores the potency CORT to inhibit uptake in the BLA (*t*8 = 2.368, *p* < .0454; Figure 7d).



7 Corticosterone inhibits uptake more in the BLA than the NAcc. (a) Representative traces from the nucleus accumbens (NAcc) at baseline and 10 µM corticosterone (CORT) overlaid. (b) Representative basolateral amygdala (BLA) traces at baseline and 10 µM CORT overlaid. (c) CORT (3 and 10 μM) significantly inhibited catecholamine uptake in the BLA compared to the NAcc, (p < .01 and p < .001, respectively). (d) CORT (10 μM) significantly increased T80 in the BLA compared to the NAcc (p < .05). NAcc, nucleus accumbens core; BLA, basolateral amygdala; CA, catecholamine; CORT, corticosterone; \*p < .05; \*\*p < .01. N = 5–8 animals/group [Colour figure can be viewed at wileyonlinelibrary.com]

## [Corticosterone increases dopamine clearance time in the NAcc of DAT KO mice](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

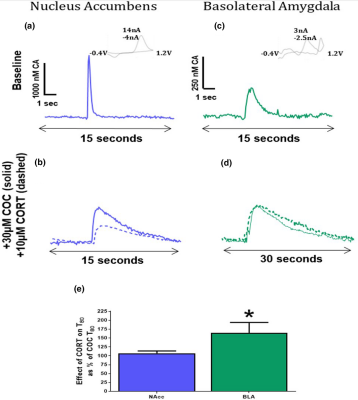
CORT treatment had a much smaller effect on inhibiting catecholamine uptake in the NAcc than in the BLA. We believed this was likely because the high levels of DAT expression in the NAcc mask the effects of CORT‐induced inhibition of OCT3. To examine the contribution of OCT3 function in the NAcc in the absence of DAT, we examined the effects of CORT on catecholamine clearance in NAcc slices obtained from DAT KO mice. A one‐way ANOVA revealed that CORT treatment increased catecholamine clearance time over the concentration response curve (*F*4,23 = 3.676; Figure 8). Tukey's post hoc analysis revealed a significant increase in T80 at the 10 µM concentration versus baseline (*p* < .05).



8 Corticosterone increases dopamine clearance time in the NAcc of DAT KO mice. Corticosterone (CORT) dose dependently decreased catecholamine clearance in DAT KO mice, indicated by an increase in T80 time (p < .05). DAT KO, dopamine transporter knockout mice; \*p < .05. N = 6 animals [Colour figure can be viewed at wileyonlinelibrary.com]

## [Catecholamine clearance in the NAcc is governed by DATs while clearance in the BLA involves O](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)CT3 activity

To better compare the relative contribution of DAT and OCT3 to catecholamine uptake in the NAcc and BLA, catecholamine uptake was measured twice in NAcc and BLA slices: first after incubation with the maximum concentration of cocaine studied here (30 µM), and second after the addition of the maximum concentration of CORT tested here (10 µM), in the presence of cocaine. Representative NAcc and BLA traces can be seen at baseline, following incubation with 30 µM cocaine, and after incubation with cocaine +10 µM CORT (Figure 9a–d). The effect of CORT on clearance was expressed as the CORT‐induced increase in T80 as a percentage of the cocaine‐induced T80. Student's *t* test revealed that magnitude of the CORT‐induced increase in T80 as a percent of cocaine‐induced changes in T80 was greater in the BLA than in the NAcc (*t*15 = 2.268, *p* < .0386; Figure 9e). Of note, it can be seen that release amplitude is reduced in the NAcc with the combination of cocaine and CORT, but this effect is not present in the BLA.



9 Catecholamine clearance in the NAcc is governed by DAT function while catecholamine clearance in the BLA favors OCT3 activity. (a) Representative traces of the nucleus accumbens (NAcc) at baseline, (b) after incubation with 30 µM cocaine (COC; solid line) and after the addition of 10 µM corticosterone (CORT; dashed line). (c) Traces from BLA slices at baseline, (d) after incubation with 30 µM COC (solid line), and after the addition of 10µM CORT (dashed line). (e) Increases in T80 due to CORT application as a percent of post‐COC clearance time revealed a greater effect of CORT on uptake inhibition in the BLA compared to the NAcc. NAcc, nucleus accumbens; BLA, basolateral amygdala; COC, cocaine (COC); CORT, corticosterone CORT; \*p < .05. N = 7–10 animals/group [Colour figure can be viewed at wileyonlinelibrary.com]

# [DISCUSSION](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

Dopaminergic signaling in the NAcc and BLA contributes to the regulation of affect (Chartoff & Carlezon, 2014; Diaz et al., 2011; Ferry et al., 2015) and motivated behaviors (Fish et al., 2014; Nakamura et al., 2014; Shiflett & Balleine, 2010; Simmons & Neill, 2009). The magnitude of catecholamine effects in a given region can be regulated by alterations in release magnitude and uptake kinetics. Previous work (Jones et al., 1995) and the present study describe reduced catecholamine release and slower uptake in the BLA compared to the NAcc. The present work aimed to expand our understanding of uptake mechanisms in these regions by examining the expression and activity of two catecholamine transporters that mediate clearance of extracellular catecholamines in the NAcc and BLA. The present studies demonstrated that the expression of DAT protein is greater in the NAcc than in the BLA and that OCT3 protein expression is similar between regions. Consistent with the transporter expression data, ex vivo voltammetry data demonstrated that the DAT inhibitor cocaine inhibits catecholamine uptake to a greater extent in the NAcc than in the BLA, while the OCT inhibitor CORT exerted a greater inhibitory effect in the BLA than in the NAcc. We suspect that differences in uptake rates are due to regional differences in the density of transporter expression. The distinct uptake mechanisms in the NAcc and BLA may contribute to differential regulation of catecholamine neurotransmission within these regions.

## [The magnitude of catecholamine release is significantly lower in the BLA than in the NAcc](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

While the present work mainly focused on differences in catecholamine uptake in the NAcc and the BLA, presynaptic mechanisms influencing catecholamine release also play a major role in shaping catecholamine signaling. The present findings using fast‐scan cyclic voltammetry corroborate previous work demonstrating that stimulated catecholamine release is significantly greater in the NAcc than in the BLA (Jones et al., 1995). While we did not further explore release mechanisms here, we and others have previously examined many potential modulators of catecholamine release that may differ between the NAcc and the BLA. Previous studies have shown that dopamine D2 and D3 autoreceptors (Maina & Mathews, 2010) and ⍺2‐adrenergic autoreceptors reduce catecholamine release (Garcia et al., 2004) in subnuclei of the NAcc (Ihalainen & Tanila, 2004; Maina & Mathews, 2010) and BLA (Ferry et al., 2015; Gurevich & Joyce, 1999; Perez de la Mora et al., 2012; Stevenson & Gratton, 2004). It is possible that differences in dopaminergic and noradrenergic autoreceptor expression between the NAcc and BLA may contribute to differential dopamine release in these two regions. Additionally, we have shown that kappa opioid receptor‐mediated inhibition of dopamine release in the NAcc is associated with negative affect‐like behaviors in rodents (Karkhanis, Rose, Weiner, & Jones, 2016; Rose et al., 2016). Kappa opioid receptors are densely expressed in both the NAcc and the BLA, where their activation leads to reductions in presynaptic release of neurotransmitters, including dopamine, and differential receptor function or expression could also influence release kinetics from these regions (Svingos, Chavkin, Colago, & Pickel, 2001; Thompson et al., 2000; Werling, Frattali, Portoghese, Takemori, & Cox, 1988). Future work aimed at differentiating release mechanisms between these regions will further aid in refining therapeutic treatment strategies for disorders of negative affect and substance use disorders.

## [Cocaine and corticosterone were more potent at inhibiting uptake rates in the NAcc and BLA, r](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)espectively

Catecholamine clearance is a complex, regionally variable process. Our Western blot and immunofluorescence findings reveal that while OCT3 is expressed at similar levels in NAcc and BLA, DAT expression is significantly greater in the NAcc than in the BLA. These data are consistent with previous studies demonstrating abundant DAT protein in the NAcc (Gurevich & Joyce, 1999; Perez de la Mora et al., 2012), and OCT3 more uniformly distributed throughout the brain (Gasser et al., 2009).

Transporter protein density is not the sole determinant of overall transporter function. Although Western blot data here show higher levels of DAT in the NAcc compared to the BLA, differences in membrane localization and transport modulators, among other factors, can contribute to differences in transporter function between regions (Kivell et al., 2014; Rao, Sorkin, & Zahniser, 2013). Studies examining the relative subcellular localizations of DAT and OCT3 and their proximity to dopamine receptors would be particularly informative. While colocalization of these transporters has not been examined (Gasser, 2018), ultrastructural studies have demonstrated that OCT3 is localized to dendritic spines and presynaptic terminals in the BLA (Gasser et al., 2017), suggesting important roles for the transporter in regulating perisynaptic monoamine concentrations. OCT3 expression has also been observed on DAT+ neurons (Mayer et al., 2018), suggesting possible co‐expression at dopamine terminals. Interestingly, OCT3 also appears to be present on striatal cholinergic neurons (Vialou, Balasse, Dumas, Giros, & Gautron, 2007), which strongly regulate terminal dynamics in the nucleus accumbens (Melchior, Ferris, Stuber, Riddle, & Jones, 2015; Siciliano, McIntosh, Jones, & Ferris, 2017; Yorgason, Rose, McIntosh, Ferris, & Jones, 2015). Therefore, it is possible that there may be additional effects of OCT3 activity on cholinergic neurons, which may alter catecholamine clearance kinetics. While localization studies will certainly be necessary for future delineation of the mechanisms of OCT3 function, we chose to focus on catecholamine clearance as a functional output of DAT and OCT3 activity.

With these studies in mind, we evaluated the functional importance of the DAT and OCT3 in the NAcc and BLA with slice voltammetry using cocaine and CORT to inhibit DAT and OCT3 function, respectively. The concentrations of both cocaine and corticosterone utilized in this study are admittedly somewhat high compared to physiological levels in the presence of systemic cocaine or stress exposure, although they are difficult to directly compare. For instance, repeated administration of 1 mg/kg cocaine over 10 injections in anesthetized rats does begin to show the characteristic decline in maximal dopamine amplitude that we observe (and have commented on here) with higher concentrations of cocaine on slice (Brodnik, Ferris, Jones, & Espana, 2017); however, this effect is not accompanied by uptake inhibition that is as dramatic as is observed with higher concentrations of cocaine on slice. Similarly, corticosterone concentrations used here are around 1–2 orders of magnitude higher than brain levels of corticosterone following administration of stressors in vivo (Makino, Gold, & Schulkin, 1994; Pitman, Ottenweller, & Natelson, 1988; Sullivan & Gratton, 1998), though the concentration response curve for corticosterone in Figure 8 does more closely approximate this range. Here, we have chosen somewhat higher concentrations that we and others have published previously in ex vivo studies in order to more clearly observe effects on reuptake (Caliskan et al., 2015; Yorgason et al., 2011).

Cocaine inhibited uptake more in the NAcc, which expresses higher levels of DAT protein, than in the BLA. The fact that CORT was more effective at reducing catecholamine uptake in the BLA than NAcc, despite our evidence that OCT3 protein levels were similar in the two regions, is likely due to the abundant expression of DAT in the NAcc. Thus, the effects of OCT3 inhibition are likely obscured by continued abundant DAT activity in the presence of CORT.

This interpretation was supported by our data examining CORT effects on catecholamine clearance in DAT KO mice, in which CORT is better able to inhibit catecholamine clearance. It is likely that CORT inhibits catecholamine uptake in the NAcc of wild‐type animals, but that the effect is not detectable due to the high levels of DAT activity. It is important to note that, as with any constitutive knockout approach, some of the effects observed in the DAT KO mouse line may be mitigated at least in part by compensatory mechanisms that could be driving altered OCT3 function in these animals. However, consistent with our findings, a recent study reported that CORT treatment decreased the clearance of naturally occurring dopamine transients in the NAcc even in the absence of DAT blockade (Wheeler et al., 2017). These studies also demonstrated that CORT pre‐treatment potentiates the effect of a previously sub‐threshold dose of cocaine on NAcc dopamine clearance.

Though the findings outlined here appear at the surface to report relatively straightforward phenomena, it should be noted that there are many factors modulating uptake kinetics and should not be viewed as overly simplistic. As noted above, the cyclic voltammograms for dopamine and norepinephrine are virtually identical, making it difficult to identify which neurotransmitter kinetics are being affected by a given treatment through voltammetry assessment alone. This is particularly problematic in the BLA, where dopamine and norepinephrine are estimated to be released at similar concentrations. Blockade of either CA either pharmacologically or using optogenetic stimulation approaches is particularly troublesome in the BLA, a region with limited stimulated catecholamine release. Though one possible solution could involve application of exogenous neurotransmitter, this approach is troublesome in that temporal kinetics observed with electrical stimulation are difficult to faithfully replicate in a slice preparation.

Other monoamine transporters in addition to DAT and OCT3 are also likely to contribute to CA clearance, especially in the BLA. The norepinephrine transporter is capable of clearing catecholamines at equal or greater efficacy than the DAT and is expressed at similar levels between the accumbens and BLA in humans (Smith, Beveridge, & Porrino, 2006). In rodents, NET plays a far larger role in catecholamine reuptake in the nucleus accumbens shell, where greater norepinephrine innervation is observed, than in the NAc core, which is primarily dopaminergic (Carboni, Silvagni, Vacca, & Di Chiara, 2006). However, in regions with high norepinephrine innervation, such as the BLA, NET consistently plays a critical role in clearance of both norepinephrine and dopamine (Carboni et al., 2006). Though beyond the scope of this study, it would be interesting to ascertain the role of NET in concert with DAT and OCT3 in differential uptake kinetics, particularly within the BLA. Additionally, other low‐affinity, high‐capacity transporters with Uptake2‐like function—OCT1, OCT2, and PMAT—may contribute to CA clearance. These transporters also function to clear monoamines from the extracellular fluid and are inhibited to CORT. Among the Uptake2 transporters, OCT3 has the highest sensitivity to CORT (Gasser, 2018) and is expressed at greater density in the NAcc and BLA than other OCT isoforms (Amphoux et al., 2006). PMAT is expressed in relatively similar levels between the NAcc and BLA (Dahlin, Xia, Kong, Hevner, & Wang, 2007) and has greater affinity for dopamine than norepinephrine (Duan & Wang, 2010). Future studies examining the role of this transporter in differential uptake kinetics between the NAcc and BLA will shed further light on Uptake2 transporters in monoamine signaling, particularly with regard to dopaminergic signaling. Interestingly, it has been recently demonstrated that CORT, via inhibition of OCT3, potentiates the effect of a low systemic dose of cocaine on dopamine clearance in the NAc (Wheeler et al., 2017). This work both differs from and compliments our own in some regards: While this study was performed in awake and behaving animals with a low dose of cocaine, we also observed further inhibition of uptake with a high concentration of cocaine over a deafferented slice when CORT was applied in the bath. It is important to note that OCT3 is canonically cocaine‐insensitive at concentrations consistent with our data (Gasser and Lowry, 2018). The ability of CORT to augment cocaine's effects of uptake inhibition resonates in behavioral studies outlined below.

Of note, we do find that CORT applied following cocaine decreases release amplitude in the NAcc, but not the BLA. While the reason for this difference is not completely clear, it is known that this combination of drugs enhances spontaneous dopamine release transients as well as reducing uptake (Graf et al., 2013; Wheeler et al., 2017), we hypothesize that elevated extracellular levels of dopamine in the slice tonically activate presynaptic D2‐type autoreceptors, which then inhibit electrically evoked release in the NAcc (Wieczorek & Kruk, 1994). The BLA may not exhibit this effect because dopamine terminals in the BLA have fewer presynaptic autoreceptors and are much less sensitive to autoreceptor‐mediated inhibition of release than in the NAcc (Garris & Wightman, 1995). These aspects of NE regulation in the BLA are unknown and require further exploration.

## [The behavioral implications of elevated corticosterone in the NAcc and BLA](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

OCT3 is directly and specifically inhibited by CORT, a hormone released in rodents during times of stress (Baganz et al., 2010; Wieland et al., 2000). Indeed, augmented levels of CORT have been measured in the NAcc (Yu et al., 2013) and BLA (Bouchez et al., 2012), following exposure to a stressor. Our findings support previous work that demonstrated CORT‐induced inhibition of dopamine clearance in the NAcc in the presence of the DAT inhibitor GBR12909 (Graf et al., 2013). These data specifically implicate OCT3 in increased extracellular catecholamine levels during times of stress. Further, through OCT3, CORT is able to augment the ability of a low dosage of systemic cocaine to inhibit uptake of dopamine in the NAcc of an awake and behaving rat (Wheeler et al., 2017). A growing body of literature suggests that augmented CORT, via inhibition of OCT3‐mediated dopamine clearance, potentiates cocaine‐induced reinstatement of drug‐seeking behavior (Graf et al., 2013; McReynolds et al., 2017). It is possible that increased CORT, and subsequent reduction in catecholamine uptake in the NAcc and BLA, drive complementary aspects of drug abuse‐related behaviors.

We hypothesize that in individuals with a substance use disorder, stress‐induced increases in glucocorticoids potentiate the effects of sub‐threshold stimuli on catecholamine concentrations in the NAcc and BLA, thus "priming" the subject to seek substances of abuse. For example, previous work has shown that increased catecholamine signaling in the BLA is anxiogenic (Bananej et al., 2012; Diaz et al., 2011), and CORT‐mediated increases in dopamine transmission may underlie stress‐induced increases in the motivation for drug self‐administration, as discussed above (Graf et al., 2013). In fact, humans (Lopez‐Castro, Hu, Papini, Ruglass, & Hien, 2015; Min, Farkas, Minnes, & Singer, 2007; Sullivan et al., 2015) and rodents (Briand & Blendy, 2013; Funk, Li, & Le, 2006) exposed to stressors have augmented drug‐seeking behaviors. Moreover, during long periods of abstinence characterized by augmented CORT levels, rats with a history of cocaine self‐administration exhibited increased drug seeking (Lack et al., 2008). Despite strong evidence supporting our hypothesis, additional in vivo work examining the specific contribution of NAcc and BLA OCT3 blockade to stress‐induced drug seeking is necessary.

# [CONCLUSIONS](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

We explored the expression of DAT and OCT3, and their contribution to catecholamine clearance, in the NAcc and BLA. Our findings demonstrate the existence of distinct mechanisms of catecholamine release and clearance in the NAcc and BLA. Specifically, we report that OCT3 blockade significantly reduces catecholamine uptake in the BLA compared to the NAcc, while cocaine is more effective at reducing uptake in the NAcc. We suspect that OCT3 removes excess synaptic catecholamines in the NAcc when transporters are saturated due to high release volumes, or when transporters are pharmacologically inhibited, for example by psychostimulant drugs of abuse. This putative DAT‐OCT3‐mediated process allows for the relative maintenance of homeostasis within the synapse under normal conditions. However, in times of stress, blockade of OCT3 is hypothesized to augment catecholamine concentrations in the BLA and NAcc and prime the subject for drug‐seeking behavior (Graf et al., 2013; McReynolds et al., 2017).

# [ACKNOWLEDGEMENTS](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

The authors would like to acknowledge Joanne Konstantopolous for her technical expertise.

# [CONFLICT OF INTERESTS](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

The authors declare no conflicts of interest.

# [AUTHOR CONTRIBUTIONS](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

KMH and JHR wrote the manuscript. JHR ran voltammetry experiments, SCF ran Western blot experiments, and JEH and KER ran immunohistochemistry experiments. PJG and SRJ edited the manuscript.

# [DATA AVAILABILITY STATEMENT](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

Data presented here can be accessed online at the following site: https://osf.io/xfqr2/?view\_only=33ecfd5e9636443891a450a964d52575.

## [Peer review](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

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## [Footnotes](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

1 Edited by Jochen Roeper

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3 [Correction added on 29 December 2020, after first online publication: Funding information details corrected.]

# [REFERENCES](https://0-web-s-ebscohost-com.libus.csd.mu.edu/ehost/detail/detail?vid=3&sid=b730cee2-9473-4208-9737-b69b8331959e%40redis&bdata=JnNpdGU9ZWhvc3QtbGl2ZSZzY29wZT1zaXRl#toc)

Akil, M., Kolachana, B. S., Rothmond, D. A., Hyde, T. M., Weinberger, D. R., & Kleinman, J. E. (2003). Catechol‐O‐methyltransferase genotype and dopamine regulation in the human brain. Journal of Neuroscience, 23 (6), 2008 – 2013.

Amphoux, A., Vialou, V., Drescher, E., Bruss, M., Mannoury La Cour, C., Rochat, C., ... Gautron, S. (2006). Differential pharmacological in vitro properties of organic cation transporters and regional distribution in rat brain. Neuropharmacology, 50 (8), 941 – 952.

Ayala‐Lopez, N., Jackson, W. F., Burnett, R., Wilson, J. N., Thompson, J. M., & Watts, S. W. (2015). Organic cation transporter 3 contributes to norepinephrine uptake into perivascular adipose tissue. American Journal of Physiology. Heart and Circulatory Physiology, 309 (11), H1904 – H1914.

Baganz, N., Horton, R., Martin, K., Holmes, A., & Daws, L. C. (2010). Repeated swim impairs serotonin clearance via a corticosterone‐sensitive mechanism: Organic cation transporter 3, the smoking gun. Journal of Neuroscience, 30 (45), 15185 – 15195.

Baik, J. H. (2013). Dopamine signaling in reward‐related behaviors. Frontiers in Neural Circuits, 7, 152.

Ballester Gonzalez, J., Dvorkin‐Gheva, A., Silva, C., Foster, J. A., & Szechtman, H. (2015). Nucleus accumbens core and pathogenesis of compulsive checking. Behavioural Pharmacology, 26 (1–2), 200 – 216.

Bananej, M., Karimi‐Sori, A., Zarrindast, M. R., & Ahmadi, S. (2012). D1 and D2 dopaminergic systems in the rat basolateral amygdala are involved in anxiogenic‐like effects induced by histamine. J Psychopharmacol, 26 (4), 564 – 574.

Bouchez, G., Millan, M. J., Rivet, J. M., Billiras, R., Boulanger, R., & Gobert, A. (2012). Quantification of extracellular levels of corticosterone in the basolateral amygdaloid complex of freely‐moving rats: A dialysis study of circadian variation and stress‐induced modulation. Brain Research, 1452, 47 – 60.

Briand, L. A., & Blendy, J. A. (2013). Not all stress is equal: CREB is not necessary for restraint stress reinstatement of cocaine‐conditioned reward. Behavioral Brain Research, 246, 63 – 68.

Brinley‐Reed, M., & McDonald, A. J. (1999). Evidence that dopaminergic axons provide a dense innervation of specific neuronal subpopulations in the rat basolateral amygdala. Brain Research, 850 (1–2), 127 – 135.

Brodnik, Z. D., Ferris, M. J., Jones, S. R., & Espana, R. A. (2017). Reinforcing doses of intravenous cocaine produce only modest dopamine uptake inhibition. ACS Chemical Neuroscience, 8 (2), 281 – 289.

Budygin, E. A., John, C. E., Mateo, Y., & Jones, S. R. (2002). Lack of cocaine effect on dopamine clearance in the core and shell of the nucleus accumbens of dopamine transporter knock‐out mice. Journal of Neuroscience, 22 (10), RC222.

Caliskan, G., Schulz, S. B., Gruber, D., Behr, J., Heinemann, U., & Gerevich, Z. (2015). Corticosterone and corticotropin‐releasing factor acutely facilitate gamma oscillations in the hippocampus in vitro. **European Journal of Neuroscience**, 41 (1), 31 – 44.

Calligaro, D. O., & Eldefrawi, M. E. (1987). High affinity stereospecific binding of [3H] cocaine in striatum and its relationship to the dopamine transporter. Membrane Biochemistry, 7 (2), 87 – 106.

Carboni, E., Silvagni, A., Vacca, C., & Di Chiara, G. (2006). Cumulative effect of norepinephrine and dopamine carrier blockade on extracellular dopamine increase in the nucleus accumbens shell, bed nucleus of stria terminalis and prefrontal cortex. Journal of Neurochemistry, 96 (2), 473 – 481.

Chartoff, E. H., Carlezon, W. A. (2014). Drug withdrawal conceptualized as a stressor. Behavioural Pharmacology, 25 (5–6), 473 – 492.

Churchwell, J. C., Morris, A. M., Heurtelou, N. M., & Kesner, R. P. (2009). Interactions between the prefrontal cortex and amygdala during delay discounting and reversal. Behavioral Neuroscience, 123 (6), 1185 – 1196.

Dahlin, A., Xia, L., Kong, W., Hevner, R., & Wang, J. (2007). Expression and immunolocalization of the plasma membrane monoamine transporter in the brain. Neuroscience, 146 (3), 1193 – 1211.

Diaz, M. R., Chappell, A. M., Christian, D. T., Anderson, N. J., & McCool, B. A. (2011). Dopamine D3‐like receptors modulate anxiety‐like behavior and regulate GABAergic transmission in the rat lateral/basolateral amygdala. Neuropsychopharmacology, 36 (5), 1090 – 1103.

Duan, H., & Wang, J. (2010). Selective transport of monoamine neurotransmitters by human plasma membrane monoamine transporter and organic cation transporter 3. Journal of Pharmacology and Experimental Therapeutics, 335 (3), 743 – 753.

Ferry, B., Parrot, S., Marien, M., Lazarus, C., Cassel, J. C., & McGaugh, J. L. (2015). Noradrenergic influences in the basolateral amygdala on inhibitory avoidance memory are mediated by an action on alpha2‐adrenoceptors. Psychoneuroendocrinology, 51, 68 – 79.

Fish, E. W., DiBerto, J. F., Krouse, M. C., Robinson, J. E., & Malanga, C. J. (2014). Different contributions of dopamine D1 and D2 receptor activity to alcohol potentiation of brain stimulation reward in C57BL/6J and DBA/2J mice. Journal of Pharmacology and Experimental Therapeutics, 350 (2), 322 – 329.

Franco, A. J., Chen, C., Scullen, T., Zsombok, A., Salahudeen, A. A., Di, S., ... Tasker, J. G. (2016). Sensitization of the hypothalamic‐pituitary‐adrenal axis in a male rat chronic stress model. Endocrinology, 157 (6), 2346 – 2355.

Funk, D., Li, Z., & Le, A. D. (2006). Effects of environmental and pharmacological stressors on c‐fos and corticotropin‐releasing factor mRNA in rat brain: Relationship to the reinstatement of alcohol seeking. Neuroscience, 138 (1), 235 – 243.

Fuxe, K., Borroto‐Escuela, D. O., Romero‐Fernandez, W., Zhang, W. B., & Agnati, L. F. Volume transmission and its different forms in the central nervous system. Chinese Journal of Integrative Medicine 2013 ; 19 (5): 323 – 329.

Gainetdinov, R. R., Jones, S. R., & Caron, M. G. (1999). Functional hyperdopaminergia in dopamine transporter knock‐out mice. Biological Psychiatry, 46 (3), 303 – 311.

Gainetdinov, R. R., Jones, S. R., Fumagalli, F., Wightman, R. M., & Caron, M. G. (1998). Re‐evaluation of the role of the dopamine transporter in dopamine system homeostasis. Brain Research Reviews, 26 (2–3), 148 – 153.

Garcia, A. S., Barrera, G., Burke, T. F., Ma, S., Hensler, J. G., & Morilak, D. A. (2004). Autoreceptor‐mediated inhibition of norepinephrine release in rat medial prefrontal cortex is maintained after chronic desipramine treatment. Journal of Neurochemistry, 91 (3), 683 – 693.

Garris, P. A., & Wightman, R. M. (1995). Distinct pharmacological regulation of evoked dopamine efflux in the amygdala and striatum of the rat in vivo. Synapse (New York, N. Y.) 20 (3): 269 – 279.

Gasser, P. J. (2018). Roles for the uptake2 transporter OCT3 in regulation of dopaminergic neurotransmission and behavior. Neurochemistry International 123 (3): 46 – 49.

Gasser, P. J., Hurley, M. M., Chan, J., & Pickel, V. M. (2017). Organic cation transporter 3 (OCT3) is localized to intracellular and surface membranes in select glial and neuronal cells within the basolateral amygdaloid complex of both rats and mice. Brain Structure and Function, 222 (4), 1913 – 1928.

Gasser, P. J., Lowry, C. A., & Orchinik, M. (2006). Corticosterone‐sensitive monoamine transport in the rat dorsomedial hypothalamus: Potential role for organic cation transporter 3 in stress‐induced modulation of monoaminergic neurotransmission. Journal of Neuroscience, 26 (34), 8758 – 8766.

Gasser, P. J., Orchinik, M., Raju, I., & Lowry, C. A. (2009). Distribution of organic cation transporter 3, a corticosterone‐sensitive monoamine transporter, in the rat brain. Journal of Comparative Neurology, 512 (4), 529 – 555.

Giros, B., Jaber, M., Jones, S. R., Wightman, R. M., & Caron, M. G. (1996). Hyperlocomotion and indifference to cocaine and amphetamine in mice lacking the dopamine transporter. Nature, 379 (6566), 606 – 612.

Graf, E. N., Wheeler, R. A., Baker, D. A., Ebben, A. L., Hill, J. E., McReynolds, J. R., ... Gasser, P. J. (2013). Corticosterone acts in the nucleus accumbens to enhance dopamine signaling and potentiate reinstatement of cocaine seeking. Journal of Neuroscience, 33 (29), 11800 – 11810.

Grundemann, D., Schechinger, B., Rappold, G. A., & Schomig, E. (1998). Molecular identification of the corticosterone‐sensitive extraneuronal catecholamine transporter. Nature Neuroscience, 1 (5), 349 – 351.

Gurevich, E. V., & Joyce, J. N. (1999). Distribution of dopamine D3 receptor expressing neurons in the human forebrain: Comparison with D2 receptor expressing neurons. Neuropsychopharmacology, 20 (1), 60 – 80.

Heien, M. L., Phillips, P. E., Stuber, G. D., Seipel, A. T., & Wightman, R. M. (2003). Overoxidation of carbon‐fiber microelectrodes enhances dopamine adsorption and increases sensitivity. Analyst, 128 (12), 1413 – 1419.

Hek, K., Direk, N., Newson, R. S., Hofman, A., Hoogendijk, W. J., Mulder, C. L., & Tiemeier, H. (2013). Anxiety disorders and salivary cortisol levels in older adults: A population‐based study. Psychoneuroendocrinology, 38 (2), 300 – 305.

Hill, J. E., & Gasser, P. J. (2013). Organic cation transporter 3 is densely expressed in the intercalated cell groups of the amygdala: Anatomical evidence for a stress hormone‐sensitive dopamine clearance system. Journal of Chemical Neuroanatomy, 52, 36 – 43.

Ihalainen, J. A., & Tanila, H. (2004). In vivo regulation of dopamine and noradrenaline release by alpha2A‐adrenoceptors in the mouse nucleus accumbens. Journal of Neurochemistry, 91 (1), 49 – 56.

Iversen, L. L., & Salt, P. J. (1970). Inhibition of catecholamine Uptake‐2 by steroids in the isolated rat heart. British Journal of Pharmacology, 40 (3), 528 – 530.

John, C. E., & Jones, S. R. (2007). Voltammetric characterization of the effect of monoamine uptake inhibitors and releasers on dopamine and serotonin uptake in mouse caudate‐putamen and substantia nigra slices. Neuropharmacology, 52 (8), 1596 – 1605.

John, C. E., & Jones, S. R. Fast scan cyclic voltammetry of dopamine and serotonin in mouse brain slices. In: A. C. Michael, & L. M. Borland, eds. Electrochemical methods for neuroscience. Boca Raton (FL) : CRC Press/Taylor & Francis, 2007.

Jones, A. B., Gupton, R., & Curtis, K. S. (2016). Estrogen and voluntary exercise interact to attenuate stress‐induced corticosterone release but not anxiety‐like behaviors in female rats. Behavioral Brain Research, 311, 279 – 286.

Jones, S. R., Garris, P. A., Kilts, C. D., & Wightman, R. M. (1995). Comparison of dopamine uptake in the basolateral amygdaloid nucleus, caudate‐putamen, and nucleus accumbens of the rat. Journal of Neurochemistry, 64 (6), 2581 – 2589.

Jung, C., Greco, S., Nguyen, H. H., Ho, J. T., Lewis, J. G., Torpy, D. J., & Inder, W. J. (2014). Plasma, salivary and urinary cortisol levels following physiological and stress doses of hydrocortisone in normal volunteers. BMC Endocrine Disorders, 14, 91.

Karkhanis, A. N., Rose, J. H., Huggins, K. N., Konstantopoulos, J. K., & Jones, S. R. (2015). Chronic intermittent ethanol exposure reduces presynaptic dopamine neurotransmission in the mouse nucleus accumbens. Drug and Alcohol Dependence, 150, 24 – 30.

Karkhanis, A. N., Rose, J. H., Weiner, J. L., & Jones, S. R. (2016). Early‐life social isolation stress increases kappa opioid receptor responsiveness and downregulates the dopamine system. Neuropsychopharmacology, 41 (9), 2263 – 2274.

Kehr, J., Hoistad, M., & Fuxe, K. (2000). Diffusion of radiolabeled dopamine, its metabolites and mannitol in the rat striatum studied by dual‐probe microdialysis. Progress in Brain Research, 125, 179 – 190.

Kivell, B., Uzelac, Z., Sundaramurthy, S., Rajamanickam, J., Ewald, A., Chefer, V., ... Shippenberg, T. S. (2014). Salvinorin A regulates dopamine transporter function via a kappa opioid receptor and ERK1/2‐dependent mechanism. Neuropharmacology, 86, 228 – 240.

Klumpers, F., Morgan, B., Terburg, D., Stein, D. J., & van Honk, J. (2015). Impaired acquisition of classically conditioned fear‐potentiated startle reflexes in humans with focal bilateral basolateral amygdala damage. Social Cognitive and Affective Neuroscience, 10 (9), 1161 – 1168.

Lack, C. M., Jones, S. R., & Roberts, D. C. (2008). Increased breakpoints on a progressive ratio schedule reinforced by IV cocaine are associated with reduced locomotor activation and reduced dopamine efflux in nucleus accumbens shell in rats. Psychopharmacology (Berl), 195 (4), 517 – 525.

Li, S. S., & McNally, G. P. (2015). A role of nucleus accumbens dopamine receptors in the nucleus accumbens core, but not shell, in fear prediction error. Behavioral Neuroscience, 129 (4), 450 – 456.

Li, Y., Zuo, Y., Yu, P., Ping, X., & Cui, C. (2015). Role of basolateral amygdala dopamine D2 receptors in impulsive choice in acute cocaine‐treated rats. Behavioral Brain Research, 287, 187 – 195.

Lightman, S. L., & Iversen, L. L. (1969). The role of uptake2 in the extraneuronal metabolism of catecholamines in the isolated rat heart. British Journal of Pharmacology, 37 (3), 638 – 649.

Lips, K. S., Volk, C., Schmitt, B. M., Pfeil, U., Arndt, P., Miska, D., ... Koepsell, H. (2005). Polyspecific cation transporters mediate luminal release of acetylcholine from bronchial epithelium. American Journal of Respiratory Cell and Molecular Biology, 33 (1), 79 – 88.

Lopez‐Castro, T., Hu, M. C., Papini, S., Ruglass, L. M., & Hien, D. A. (2015). Pathways to change: Use trajectories following trauma‐informed treatment of women with co‐occurring post‐traumatic stress disorder and substance use disorders. Drug and Alcohol Review, 34 (3), 242 – 251.

Maina, F. K., & Mathews, T. A. (2010). A functional fast scan cyclic voltammetry assay to characterize dopamine D2 and D3 autoreceptors in the mouse striatum. ACS Chemical Neuroscience, 1 (6), 450 – 462.

Makino, S., Gold, P. W., & Schulkin, J. (1994). Corticosterone effects on corticotropin‐releasing hormone mRNA in the central nucleus of the amygdala and the parvocellular region of the paraventricular nucleus of the hypothalamus. Brain Research, 640 (1–2), 105 – 112.

Markou, A., & Koob, G. F. (1992). Bromocriptine reverses the elevation in intracranial self‐stimulation thresholds observed in a rat model of cocaine withdrawal. Neuropsychopharmacology, 7 (3), 213 – 224.

Mayer, F. P., Schmid, D., Owens, W. A., Gould, G. G., Apuschkin, M., Kudlacek, O., ... Sitte, H. H. (2018). An unsuspected role for organic cation transporter 3 in the actions of amphetamine. Neuropsychopharmacology, 43 (12), 2408 – 2417.

McReynolds, J. R., Taylor, A., Vranjkovic, O., Ambrosius, T., Derricks, O., Nino, B., ... Mantsch, J. R. (2017). Corticosterone potentiation of cocaine‐induced reinstatement of conditioned place preference in mice is mediated by blockade of the organic cation transporter 3. Neuropsychopharmacology, 42 (3), 757 – 765.

Melchior, J. R., Ferris, M. J., Stuber, G. D., Riddle, D. R., & Jones, S. R. (2015). Optogenetic versus electrical stimulation of dopamine terminals in the nucleus accumbens reveals local modulation of presynaptic release. Journal of Neurochemistry, 134 (5), 833 – 844.

Min, M., Farkas, K., Minnes, S., & Singer, L. T. (2007). Impact of childhood abuse and neglect on substance abuse and psychological distress in adulthood. Journal of Traumatic Stress, 20 (5), 833 – 844.

Nakamura, T., Sato, A., Kitsukawa, T., Momiyama, T., Yamamori, T., & Sasaoka, T. (2014). Distinct motor impairments of dopamine D1 and D2 receptor knockout mice revealed by three types of motor behavior. Frontiers in Integrative Neuroscience, 8, 56.

Napolitano, A., Bellini, G., Borroni, E., Zurcher, G., & Bonuccelli, U. (2003). Effects of peripheral and central catechol‐O‐methyltransferase inhibition on striatal extracellular levels of dopamine: A microdialysis study in freely moving rats. Parkinsonism & Related Disorders, 9 (3), 145 – 150.

Ottenweller, J. E., Natelson, B. H., Pitman, D. L., & Drastal, S. D. (1989). Adrenocortical and behavioral responses to repeated stressors: Toward an animal model of chronic stress and stress‐related mental illness. Biological Psychiatry, 26 (8), 829 – 841.

Perez de la Mora, M., Gallegos‐Cari, A., Crespo‐Ramirez, M., Marcellino, D., Hansson, A. C., & Fuxe, K. (2012). Distribution of dopamine D(2)‐like receptors in the rat amygdala and their role in the modulation of unconditioned fear and anxiety. Neuroscience, 201, 252 – 266.

Perez de la Mora, M., Jacobsen, K. X., Crespo‐Ramirez, M., Flores‐Gracia, C., & Fuxe, K. (2008). Wiring and volume transmission in rat amygdala. Implications for fear and anxiety. Neurochemical Research, 33 (8), 1618 – 1633.

Pitman, D. L., Ottenweller, J. E., & Natelson, B. H. (1988). Plasma corticosterone levels during repeated presentation of two intensities of restraint stress: Chronic stress and habituation. Physiology & Behavior, 43 (1), 47 – 55.

Rao, A., Sorkin, A., & Zahniser, N. R. (2013). Mice expressing markedly reduced striatal dopamine transporters exhibit increased locomotor activity, dopamine uptake turnover rate, and cocaine responsiveness. Synapse (New York, N. Y.), 67 (10), 668 – 677.

Rose, J. H., Karkhanis, A. N., Chen, R., Gioia, D., Lopez, M. F., Becker, H. C., ... Jones, S. R. (2016). Supersensitive kappa opioid receptors promotes ethanol withdrawal‐related behaviors and reduce dopamine signaling in the nucleus accumbens. International Journal of Neuropsychopharmacology, 19 (5), pyv127.

Schmitt, A., Mossner, R., Gossmann, A., Fischer, I. G., Gorboulev, V., Murphy, D. L., ... Lesch, K. P. (2003). Organic cation transporter capable of transporting serotonin is up‐regulated in serotonin transporter‐deficient mice. Journal of Neuroscience Research, 71 (5), 701 – 709.

Schulteis, G., Markou, A., Cole, M., & Koob, G. F. (1995). Decreased brain reward produced by ethanol withdrawal. Proceedings of the National Academy of Sciences, USA, 92 (13), 5880 – 5884.

Shiflett, M. W., & Balleine, B. W. (2010). At the limbic‐motor interface: Disconnection of basolateral amygdala from nucleus accumbens core and shell reveals dissociable components of incentive motivation. **European Journal of Neuroscience**, 32 (10), 1735 – 1743.

Siciliano, C. A., Calipari, E. S., Ferris, M. J., & Jones, S. R. (2015). Adaptations of presynaptic dopamine terminals induced by psychostimulant self‐administration. ACS Chemical Neuroscience, 6 (1), 27 – 36.

Siciliano, C. A., Calipari, E. S., Yorgason, J. T., Lovinger, D. M., Mateo, Y., Jimenez, V. A., ... Jones, S. R. (2016). Increased presynaptic regulation of dopamine neurotransmission in the nucleus accumbens core following chronic ethanol self‐administration in female macaques. Psychopharmacology (Berl), 233 (8), 1435 – 1443.

Siciliano, C. A., McIntosh, J. M., Jones, S. R., & Ferris, M. J. (2017). alpha6beta2 subunit containing nicotinic acetylcholine receptors exert opposing actions on rapid dopamine signaling in the nucleus accumbens of rats with high‐versus low‐response to novelty. Neuropharmacology, 126, 281 – 291.

Simmons, D. A., & Neill, D. B. (2009). Functional interaction between the basolateral amygdala and the nucleus accumbens underlies incentive motivation for food reward on a fixed ratio schedule. Neuroscience, 159 (4), 1264 – 1273.

Smith, H. R., Beveridge, T. J., & Porrino, L. J. (2006). Distribution of norepinephrine transporters in the non‐human primate brain. Neuroscience, 138 (2), 703 – 714.

Smyth, N., Thorn, L., Oskis, A., Hucklebridge, F., Evans, P., & Clow, A. (2015). Anxious attachment style predicts an enhanced cortisol response to group psychosocial stress. Stress, 18 (2), 143 – 148.

Stevenson, C. W., & Gratton, A. (2004). Basolateral amygdala dopamine receptor antagonism modulates initial reactivity to but not habituation of the acoustic startle response. Behavioral Brain Research, 153 (2), 383 – 387.

Sullivan, R. M., & Gratton, A. (1998). Relationships between stress‐induced increases in medial prefrontal cortical dopamine and plasma corticosterone levels in rats: Role of cerebral laterality. Neuroscience, 83 (1), 81 – 91.

Sullivan, T. P., Flanagan, J. C., Dudley, D. N., Holt, L. J., Mazure, C. M., & McKee, S. A. (2015). Correlates of smoking status among women experiencing intimate partner violence: Substance use, posttraumatic stress, and coping. American Journal on Addictions, 24 (6), 546 – 553.

Svingos, A. L., Chavkin, C., Colago, E. E., & Pickel, V. M. (2001). Major coexpression of kappa‐opioid receptors and the dopamine transporter in nucleus accumbens axonal profiles. Synapse (New York, N. Y.), 42 (3), 185 – 192.

Thompson, A. C., Zapata, A., Justice, J. B., Vaughan, R. A., Sharpe, L. G., & Shippenberg, T. S. (2000). Kappa‐opioid receptor activation modifies dopamine uptake in the nucleus accumbens and opposes the effects of cocaine. Journal of Neuroscience, 20 (24), 9333 – 9340.

Touzani, K., Bodnar, R., & Sclafani, A. (2008). Activation of dopamine D1‐like receptors in nucleus accumbens is critical for the acquisition, but not the expression, of nutrient‐conditioned flavor preferences in rats. **European Journal of Neuroscience**, 27 (6), 1525 – 1533.

Trouillon, R., Lin, Y., Mellander, L. J., Keighron, J. D., & Ewing, A. G. (2013). Evaluating the diffusion coefficient of dopamine at the cell surface during amperometric detection: Disk vs ring microelectrodes. Analytical Chemistry, 85 (13), 6421 – 6428.

Uhl, G. R., Hall, F. S., & Sora, I. (2002). Cocaine, reward, movement and monoamine transporters. Molecular Psychiatry, 7 (1), 21 – 26.

Vialou, V., Amphoux, A., Zwart, R., Giros, B., & Gautron, S. (2004). Organic cation transporter 3 (Slc22a3) is implicated in salt‐intake regulation. Journal of Neuroscience, 24 (11), 2846 – 2851.

Vialou, V., Balasse, L., Dumas, S., Giros, B., & Gautron, S. (2007). Neurochemical characterization of pathways expressing plasma membrane monoamine transporter in the rat brain. Neuroscience, 144 (2), 616 – 622.

Werling, L. L., Frattali, A., Portoghese, P. S., Takemori, A. E., & Cox, B. M. (1988). Kappa receptor regulation of dopamine release from striatum and cortex of rats and guinea pigs. Journal of Pharmacology and Experimental Therapeutics, 246 (1), 282 – 286.

Wheeler, D. S., Ebben, A. L., Kurtoglu, B., Lovell, M. E., Bohn, A. T., Jasek, I. A., ... Wheeler, R. A. (2017). Corticosterone regulates both naturally occurring and cocaine‐induced dopamine signaling by selectively decreasing dopamine uptake. **European Journal of Neuroscience**, 46 (10), 2638 – 2646.

Wieczorek, W. J., & Kruk, Z. L. (1994). A quantitative comparison on the effects of benztropine, cocaine and nomifensine on electrically evoked dopamine overflow and rate of re‐uptake in the caudate putamen and nucleus accumbens in the rat brain slice. Brain Research, 657 (1–2), 42 – 50.

Wieland, A., Hayer‐Zillgen, M., Bonisch, H., & Bruss, M. (2000). Analysis of the gene structure of the human (SLC22A3) and murine (Slc22a3) extraneuronal monoamine transporter. Journal of Neural Transmission (Vienna), 107 (10), 1149 – 1157.

Yetnikoff, L., Lavezzi, H. N., Reichard, R. A., & Zahm, D. S. (2014). An update on the connections of the ventral mesencephalic dopaminergic complex. Neuroscience, 282, 23 – 48.

Yorgason, J. T., Calipari, E. S., Ferris, M. J., Karkhanis, A. N., Fordahl, S. C., Weiner, J. L., & Jones, S. R. (2016). Social isolation rearing increases dopamine uptake and psychostimulant potency in the striatum. Neuropharmacology, 101, 471 – 479.

Yorgason, J. T., Espana, R. A., & Jones, S. R. (2011). Demon voltammetry and analysis software: Analysis of cocaine‐induced alterations in dopamine signaling using multiple kinetic measures. Journal of Neuroscience Methods, 202 (2), 158 – 164.

Yorgason, J. T., Rose, J. H., McIntosh, J. M., Ferris, M. J., & Jones, S. R. (2015). Greater ethanol inhibition of presynaptic dopamine release in C57BL/6J than DBA/2J mice: Role of nicotinic acetylcholine receptors. Neuroscience, 284, 854 – 864.

Yu, P., An, S., Tai, F., Wang, J., Wu, R., & Wang, B. (2013). Early social deprivation impairs pair bonding and alters serum corticosterone and the NAcc dopamine system in mandarin voles. Psychoneuroendocrinology, 38 (12), 3128 – 3138.

Zhang, J., Muller, J. F., & McDonald, A. J. (2013). Noradrenergic innervation of pyramidal cells in the rat basolateral amygdala. Neuroscience, 228, 395 – 408.

Zhu, H. J., Appel, D. I., Grundemann, D., & Markowitz, J. S. (2010). Interaction of organic cation transporter 3 (SLC22A3) and amphetamine. Journal of Neurochemistry, 114 (1), 142 –149.