The role of interindividual licking received and dopamine genotype on later-life licking provisioning in female rat offspring

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\textbf{Abstract}

\textbf{Introduction:} Rat mothers exhibit natural variations in care that propagate between generations of female offspring. However, there is limited information on genetic variation that could influence this propagation.

\textbf{Methods:} We assessed early-life maternal care received by individual female rat offspring, later-life maternal care provisioning, and dopaminergic activity in the maternal brain in relation to naturally occurring genetic polymorphisms linked to the dopaminergic system. We also conducted a systematic analysis of other genetic variants potentially related to maternal behavior in our Long-Evans rat population.

\textbf{Results:} While we did not find a direct relationship between early-life licking received and later-life licking provisioning, this relationship was indirectly affected by dopamine levels in the nucleus accumbens and dependent on variation in the dopamine receptor 2 gene (rs107017253). More specifically, female rat offspring with the A/G genotype showed a positive relationship between average licking received and dopamine levels in the nucleus accumbens of the maternal brain; there was no relationship with female rat offspring with the A/A genotype. The higher dopamine levels in the nucleus accumbens corresponded with higher maternal licking provisioning from postnatal days 2–9. We also discovered and validated several new variants that were predicted by our systematic analysis.

\textbf{Conclusion:} Our findings suggest that genetic variation influences the relationship between early-life maternal care received and the dopaminergic system of the maternal brain, which can indirectly influence later-life maternal care provisioning.

\textbf{Keywords} dopamine receptor D2, female, gene × environment interaction, genotype, individual differences, intergenerational, long-evans, maternal care, moderated mediation, rat
INTRODUCTION

The maternal environment has a substantial role in influencing offspring phenotype in adulthood, including the intergenerational transmission of maternal care through the female lineage (Fleming et al., 2002). Both maternal care received and changes in the brain that occur when the mother is preparing for postnatal offspring care involve alterations in the dopaminergic (Champagne et al., 2004; Peña et al., 2014) and oxytocinergic systems (Beery et al., 2016; Francis et al., 2002; Toepfer et al., 2017), as well as changes in estrogen signaling (Cameron et al., 2008; Champagne, 2008). Maternal care provisioning also involves neuropeptides aside from oxytocin, including prolactin and arginine vasopressin (Bridges, 2015). These mechanisms have been well-characterized in rodents, and studies with humans show similar findings (Rosenblatt, 1994), suggesting that the factors involved in the intergenerational transmission of maternal care are evolutionarily conserved.

Rat pups with early-life adversity induced by maternal and sibling deprivation are less attentive to foster pups and their own offspring later in life than rat pups maternally reared (Gonzalez et al., 2001; Palombo et al., 2010). Higher licking-like tactile stimulation during the deprivation period partially mitigates the later-life impairments in maternal care provisioning (Gonzalez et al., 2001; Palombo et al., 2010). In addition, naturally occurring variation between litters in maternal licking/grooming received can transmit between generations of offspring. High licking/grooming and low licking/grooming mothers (defined as ± one standard deviation from the average) tend to produce female offspring that become high licking/grooming and low licking/grooming mothers to their offspring, respectively (Champagne et al., 2003; Francis et al., 1999). This effect persists in cross-fostering experiments, suggesting a prominent role of maternal behavior in the intergenerational transmission of maternal care (Francis et al., 1999).

Previous work on the intergenerational transmission of maternal care in rats has mainly focused on proxies of maternal neglect or the tail ends of the normal distribution of maternal care received. We have more limited knowledge of factors that affect the intergenerational transmission of maternal care in the average rat mother. Seminal research examining natural variations in maternal care received between litters found that there is also substantial within-litter variation in later-life maternal care provisioning (Champagne et al., 2003). In addition, studies with human cohorts in nonclinical populations have shown a modest relationship between parental care received and parental care provisioning (Van Ijzendoorn, 1992) and found that the genotype of the individual can interact with the early-life environment or directly affect maternal care provisioning (Fleming & Kraemer, 2019). Studies in mice also show an important role of offspring genotype in maternal care received (Ashbrook et al., 2015, 2017).

Findings from our group and others have shown effects of early-life interindividual maternal care received on later-life behavior (Hasselt et al., 2012; Lauby et al., 2019; Pan et al., 2014, 2018; Ragan et al., 2012). We have also reported gene x environment interactions involved in some of these effects (Lauby et al., 2019; Pan et al., 2018). For example, we recently reported an interaction between a dopamine receptor 2 (Drd2) single nucleotide polymorphism (rs107017253) and early-life average licking duration per bout (a proxy of maternal care quality) on strategy shifting and sucrose preference, two dopamine-related phenotypes (Lauby et al., 2019). Executive function measured by strategy shifting performance has previously been shown to be related to maternal care provisioning in human cohorts (Gonzalez et al., 2012). However, studies on the effect of genotype in the Long-Evans rat model commonly used in these experiments have been limited by the lack of knowledge of genetic variation in this outbred strain. Although there are now data available on genetic variation within wild rat populations (Ness et al., 2012) and between inbred rat lines (Hermsen et al., 2015; STAR Consortium et al., 2008), no mapping has been carried out within the Long-Evans population.

The purpose of this study was to investigate the role of genotype and gene x environment interactions in the relationship between early-life maternal licking received and later-life licking provisioning.

![Diagram](https://via.placeholder.com/150)

**FIGURE 1** Hypothesized moderated mediation between early-life licking received and later-life licking provisioning. Interindividual licking received early in life would positively associate with later-life maternal licking provisioning. Differences in dopaminergic activity in the maternal brain would account for this association and mediate the relationship between licking received and licking provisioning indirectly. Finally, offspring genotype in dopamine-related genes would interact with early-life interindividual licking received and moderate later-life maternal licking provisioning and dopaminergic activity in the maternal brain.
with a specific focus on the dopaminergic system of the maternal brain. We also aimed to investigate other genes relevant to maternal behavior using a more systematic approach to identify and verify genetic variants in our Long-Evans rat population. We measured interindividual maternal licking received within the first week of life and assessed later-life female offspring maternal licking provisioning, maternal brain dopamine (DA) and its metabolite levels, and variation in single nucleotide polymorphisms (SNPs) in genes involved in DA, estrogen, and neuropeptide function. We predicted that higher early-life licking received would correspond to higher later-life licking provisioning in female offspring. In addition, we hypothesized that this relationship would be indirectly affected by dopaminergic activity in the maternal brain and interact with female offspring genotype. The hypothesized moderated mediation model is displayed in Figure 1.

2 | MATERIALS AND METHODS

2.1 | Rat breeding

Seven-week-old female \( n = 24 \) and male \( n = 6 \) Long-Evans rats were obtained from Charles River Laboratories. They were housed in same-sex pairs on a 12:12-hr light–dark cycle (lights on at 7:00) with ad libitum access to standard chow diet and water. For breeding, one male was housed with two females for 1 week. Females were then housed separately and weighed weekly throughout pregnancy. The breeding males were used multiple times to produce four cohorts of litters in this study. All animal procedures were approved by the Local Animal Care Committee at the University of Toronto Scarborough and conformed to the guidelines of the Canadian Council on Animal Care.

The pregnant \( F_0 \) females were checked for parturition starting three weeks after breeding at 9:00 and 17:00. Postnatal day (PND) 0 was determined whether the birth occurred between 9:00 and 17:00 or if pups were noted at 9:00 but have not nursed yet. Pups found at 9:00 with a visible milk band were determined to be PND 1. At PND 1, litters were culled to five to six female pups and individually weighed. We focused on smaller litters in order to accurately measure maternal care received. Therefore, male offspring were not examined for this study. A total of 136 \( F_1 \) female pups were assessed for maternal care received.

2.2 | Maternal care received observations

At PND 1, 3, 5, and 7, maternal care was assessed as previously reported (Lauby et al., 2019). From 10:00 to 17:00 in the light phase, litters were briefly separated from their mother (for 10 to 20 min) and individually marked using odorless and tasteless food coloring (Club House, London, Canada) to distinguish between siblings. The entire litter was then placed in the opposite corner of the established nest, and maternal behavior was observed for 30 min using a Leica CM3050S cryostat (Leica Microsystems, Wetzlar, Germany). Observer XT 11.5 (Noldus Information Technology). To establish inter-rater reliabilities on behavioral observations, three researchers coded the same mothers with an experienced coder until high reliability (>90%) was consistently met. Cages were not changed throughout the maternal behavior observation period.

Duration and frequency of anogenital licking and body licking were coded for individual pups, meaning each pup had designated keys in the Observer software. Total duration of licking (anogenital and body licking) and average duration of a lick bout (total duration of licking/number of licking bouts) across all four observation days for each pup were calculated as measures of maternal care. This was referred to as “total licking duration" and “average licking duration," respectively.

Female offspring were weaned at PND 22 and were pair-housed with siblings. They were weighed periodically until adulthood (PND 75).

2.3 | Maternal care provisioning observations

A subset of adult \( F_1 \) female offspring \( n = 54 \) were bred with sexually experienced males \( n = 6 \) for one week. Six females were unable to get pregnant after two rounds of breeding and two females failed to lactate following parturition, with a total of 46 \( F_1 \) rat mothers being used for intergenerational maternal care analysis. Births were checked starting three weeks after breeding at 9:00 and 17:00. At PND 1, cages were changed with no culling of pups. The litter sizes ranged from 4 to 18 pups and were not significantly correlated with maternal licking provisioning (Pearson’s \( r = 0.001 \), \( p = .994 \)).

From postnatal days 2–9, each litter was video recorded for one hour three times during the light phase (9:00–10:00, 13:00–14:00, 17:00–18:00) and three times during the dark phase (21:00–22:22, 1:00–2:00, 5:00–6:00) using security cameras connected to a DVR system (Swann Communications Ltd.). These videos were coded with Observer XT 10.5 (Noldus Information Technology) for maternal behavior by four coders who were blind to the genotype of the \( F_1 \) rat mother. Nursing, licking, nest-building, and other self-directed behaviors were scored every three minutes based on previous literature (Champagne et al., 2003). A total of 120 observations per \( F_1 \) mother per day were coded, and maternal licking provisioning was represented as a percentage of the frequency of licking behavior (body and anogenital combined) coded over total observations.

After 10:00 on PND 9 of the \( F_2 \) pups, the \( F_1 \) rat mother was separated from her pups and was sacrificed with \( CO_2 \) inhalation and decapitation. Liver and whole brain tissue were collected and placed in dry ice or flash-frozen in isopentane, respectively. Tissue was stored in −80°C until further processing.

2.4 | High-performance liquid chromatography

Forty-six maternal rat brains were sliced and microdissected using
Medial prefrontal cortex (+4.20 mm to +2.70 mm Bregma), nucleus accumbens core and shell (+2.20 mm to +1.20 mm Bregma), medial preoptic area (0.30 mm to −0.80 mm Bregma), dorsal hippocampus (control brain region; −2.30 mm to −3.30 mm Bregma), and the ventral tegmental area (−5.20 mm to −5.60 mm Bregma) were identified using an adult rat brain atlas (Paxinos & Watson, 1997). 5 μl of 1.0 M ascorbic acid was added to each sample to stabilize the neurotransmitters followed by storage at −80°C.

To prepare the samples, the brain tissue was thawed on ice, suspended in 20 μl artificial cerebrospinal fluid (Harvard Apparatus, Holliston, MA), and homogenized by four pulses of sonication (2 together with a Uniget C-18 reverse phase microbore column (#8949, 4°C. The supernatant was collected and stored at −80°C. Remaining homogenate and was centrifuged at 9,600 rpm. The last 1 μl of 0.2 M perchloric acid per sample was added to the remaining homogenate and was centrifuged at 9,600 g for 10 min at 4°C. The supernatant was collected and stored at −80°C.

Dopamine (DA) and 3, 4-dihydroxyphenylacetic acid (DOPAC) levels were measured by high-performance liquid chromatography (HPLC) as described by previous studies (Chatterjee & Gerlai, 2009; Lauby et al., 2019). A BAS 460 MICROBORE-HPLC system with electrochemical detection (Bio-analytical Systems Inc.) was used together with a Uniget C-18 reverse phase microbore column (#8949, Bio-analytical Systems Inc.). The mobile phase contained a buffer [0.1 M monochloro acetic acid, 0.5 mM Na-EDTA, 0.15 g/L Na- octylsulfonate, and 10 mM sodium chloride, pH 3.1], acetonitrile, and tetrahydrofuran at a ratio of 94:3.5:0.7. The flow rate was 1.0 ml/min, and the working electrode (Uniget 3 mm glassy carbon, BAS P/N MF-1003) was set at 550 mV versus. Ag/Ag/Cl reference electrode. Detection gain was 1.0 nA, filter was 0.2 Hz, and detection limit was set at 20 nA. 5 μl of the sample supernatant was directly injected into the column. External standards for DA and DOPAC (Sigma Aldrich) of known concentrations were used to quantify and identify peaks on the chromatogram. Under these parameters, the retention times for DA and DOPAC were approximately 3.7 and 5.5 min, respectively.

DA and 3, 4-dihydroxyphenylacetic acid (DOPAC) levels were normalized against total protein concentration for each sample. DOPAC divided by DA, or the DOPAC/DA ratio, was calculated as a measure of dopaminergic activity in each brain area. We were unable to analyze two prefrontal cortex samples, two nucleus accumbens samples, one medial preoptic area sample, and three ventral tegmental area samples due to lost tissue during collection.

2.5 | Genotyping

To identify variants which segregate within the Long-Evans population, we took advantage of large amounts of RNA-seq data available online in the population. We identified 119 RNA-seq datasets on the Gene Expression Omnibus (GEO; Table S1) which we could use for variant calling. Fastq files were downloaded, aligned and variants called following the Genome Analysis Toolkit (GATK) guidelines. In brief, adaptors were trimmed from the fastq files using trim galore (version 0.4.1). The aligned files were aligned to the m6 rat reference genome using STAR (version 2.6.0), using two-pass mode (Dobin et al., 2013). Read groups were added, and duplicates were marked using picard tools. NCigar reads were split using GATK (version 4.0.8.1), and variants were called using HaplotypeCaller. Joint calling was carried out using all 119 samples using GenomicsDBImport and GenotypeGVCFs. Variant recalibration was carried out, using a “true-positive” variant list from the rat genome database. This provided us with a list of high-quality single nucleotide polymorphisms (SNPs) and small INDELs.

To validate variation in candidate SNPs, liver DNA from the F1 rat mothers was extracted using an EZNA Tissue DNA extraction kit (Omega Bio-Tek) and assessed for SNPs in genes relevant to maternal behavior, broadly including DA-related, estrogen-related, and neuropeptide-related genes (Table 1). Purified DNA (35 ng/μl) was submitted for a multiplex assay at The Centre for Applied Genomics ( SickKids, Toronto, Canada).

2.6 | Statistical analysis

All statistical analyses were performed using SPSS (IBM Corporation). We removed two extreme outliers (>3 SD from the mean) from the nPFC DOPAC/DA ratio dataset and three extreme outliers from the MPOA DOPAC/DA ratio dataset. To examine the relationship between early-life maternal licking received and later-life maternal licking provisioning, a Pearson correlation was used between total or average licking duration received and the percentage of licking provisioning observed. In addition, to examine the relationship between later-life maternal licking provisioning and dopaminergic activity, a Pearson correlation was used between percentage of licking provisioning observed and the DOPAC/DA ratio in each brain area, with a false discovery rate (FDR) correction using the Benjamini-Hochberg procedure to account for multiple analyses. Significant correlations with DOPAC/DA ratio in a brain area were followed up with Pearson’s correlations with DOPAC and DA levels separately. To examine the effects of genotype, a one-way ANOVA was used to compare offspring with each varying genotype to early-life maternal licking received and later-life maternal licking provisioning with a FDR correction to account for multiple SNP analyses for each outcome licking measure. Significant effects of genotype were followed with a Tukey’s post hoc test. To examine gene x environment interactions and the mediating role of dopaminergic activity on later-life maternal licking provisioning, we used Hayes PROCESS module for SPSS (Version 3.2) using a simple moderation (Model 1) with a FDR correction to account for multiple analyses for each outcome-dependent variable and a moderated mediation (Model 7) (Hayes, 2013). A moderation tests for an interaction between an independent variable (X) and a moderator variable (W) on a dependent variable (Y) and a mediation tests for indirect associations of the independent (X) and dependent variable (Y) by a third causal variable (the mediator M). PROCESS is a flexible modeling module that can conduct moderation and mediation analyses using multiple regression and conduct post hoc analyses of the conditional effects of a focal moderator and the indirect effects of a mediator.
The statistical model used for a simple moderation (Model 1) was as follows:

- Conditional direct effects of $X$ on $Y = b_1 + b_3W$

The statistical model used for a moderated mediation (Model 7) was as follows:

- Conditional indirect effects of $X$ on $Y$ thorough $M_i = (a_{1i} + a_{3i}W)b_i$
- Direct effect of $X$ on $Y = c'$

Multicategorical moderator variables (i.e., more than two genotypes) were analyzed using the indicator coding system. All effects were considered statistically significant if $p \leq 0.05$ and marginally significant if $p \leq 0.10$.

3 | RESULTS

3.1 | Relationship between early-life licking received and later-life licking provisioning

Total licking duration received across all maternal care observation periods (PND 1, 3, 5, and 7) averaged 168.79 ± 18.6 s and ranged from 0 to 450.72 s (Figure 2a). Average licking duration per bout of licking received averaged 8.02 ± 0.56 s and ranged from 0 to 13.93 s (Figure 2b).

We first examined the relationship between early-life maternal care received and later-life maternal care provisioning. There were no direct associations between early-life licking received and later-life licking provisioning. Neither total licking received (Pearson's $r = -0.036, p = .812$; Figure 2c) nor average licking received (Pearson's $r = -0.027, p = .861$; Figure 2d) significantly correlated with later-life licking provisioning.

3.2 | Relationship between early-life licking received, later-life licking provisioning, and dopaminergic activity in the maternal brain

We then examined relationships between maternal care received and provisioning with DOPAC/DA ratio in the medial prefrontal cortex, nucleus accumbens, medial preoptic area, and ventral tegmental area in the F1 maternal brain. There was a significant negative correlation between later-life licking provisioning and DOPAC/DA ratio in the nucleus accumbens (Pearson's $r = -0.389$, FDR adjusted $p = .036$; Figure 3a). Upon further analysis of DOPAC and DA levels separately, there was a significant positive correlation between later-life licking provisioning and DA levels (Pearson's $r = 0.346$, $p = .021$; Figure 3b) and no correlation with DOPAC levels (Pearson's

### TABLE 1 List of candidate single nucleotide polymorphisms assessed for variation, information related to their location and function, and genotype frequencies in our Long-Evans rat population

| Gene                  | Coordinate | Location | Consequence | Accession number | Genotype frequencies |
|-----------------------|------------|----------|-------------|------------------|----------------------|
| Dopamine-Related      |            |          |             |                  |                      |
| Dopamine Receptor 1 (Drd1) | chr17:11100256 | Upstream | Unknown | rs105271298 | 33/T/T, 12/T/A, 1 A/A |
| Dopamine Receptor 2 (Drd2) | chr8:53736747 | Exon | Synonymous | N/A | No variation |
| Dopamine Transporter (Dat) | chr1:32354790 | Exon | Synonymous | rs13448119 | 12 A/A, 26 A/G, 8 G/G*** |
| |               | Exon | Synonymous | N/A | 12 A/A, 26 A/G, 8 G/G*** |
| |               | Exon | Synonymous | rs13448119 | 13 G/G, 25 G/A, 8 A/A*** |
| Estrogen-Related      |            |          |             |                  |                      |
| Estrogen Receptor Alpha (Esr1) | chr1:41325782 | Exon | Synonymous | N/A | 15 C/C, 31 C/T |
| Estrogen Receptor Beta (Esr2) | chr6:99180312 | Exon | Synonymous | rs8170138 | 37 A/A, 9 A/G*** |
| Aromatase (Cyp19a1)   | chr8:58755644 | Exon | Synonymous | rs10679862 | 37 C/C, 9 C/T*** |
| |               | Exon | Missense | rs197770329 | No variation |
| |               | Exon | Synonymous | N/A | 13 T/T, 21 T/C, 12 C/C |
| Neuropeptide-Related  |            |          |             |                  |                      |
| Prolactin Receptor (Prlr) | chr2:60285990 | Exon | Synonymous | rs199216170 | No variation |
| Arginine Vasopressin (Avp) | chr3:123118892 | Intron | Unknown | rs197835086 | No variation |
| Arginine Vasopressin Receptor 1a (Avpr1a) | chr7:67341790 | Intron | Unknown | rs199089038 | No variation |
| Oxytocin Receptor (Oxtr) | chr4:144398803 | Downstream | Unknown | N/A | 23 A/A, 23 G/G |
| *,**SNPs are in linkage disequilibrium.
r = -0.060, p = .701; Figure 3c) in the nucleus accumbens. There were no other associations between later-life licking provisioning and DOPAC/DA ratio in any other brain regions.

3.3 | Effects of genotype

We detected 792,253 SNPs which segregate in the Long-Evans population. These variants are available as a UCSC genome browser track at http://genome.ucsc.edu/cgi-bin/hgTracks?db=mm6&lastViewModeType=default&hgsid=939469837_sHR5zER8w8j4UfQ58Yl6PaLa. We chose seven genes relevant to maternal care provisioning and examined 16 variants within these genes.

Of the 16 variants in genes relevant to maternal care provisioning detected, 11 showed variation in our animals (Table 1). These SNPs were in dopamine receptor 1 (Drd1; rs105271298), dopamine receptor 2 (Drd2; rs107017253), dopamine transporter (Dat; chr1:32354790 and rs13448119), estrogen receptor alpha (Esr1; chr1:41325782 and chr1:41590586), estrogen receptor beta (Esr2; rs8170138, rs13448055, and rs106738562), aromatase (Cyp19a1; chr8:58755644), and oxytocin receptor (Oxtr; chr4:144398803). We did not detect variation in the SNPs we tested in prolactin receptor (Prlr), arginine vasopressin (Avp), and arginine vasopressin receptor 1a (Avpr1a).

The one female offspring with the Drd1 A/A genotype was removed from subsequent analyses of Drd1 genotype. In addition, since the three SNPs within Esr2 and the two SNPs within Dat were in complete linkage, only one SNP in each gene (rs8170138 and rs13448119, respectively) was used for analysis.

There was a main effect of Oxtr (chr4:144398803) genotype ($F_{1,44} = 9.14$, FDR adjusted $p = 0.032$) and a marginal effect of Esr2 (rs8170138) genotype ($F_{1,44} = 6.78$, FDR adjusted $p = 0.056$) and Cyp19a1 (chr8:58755644) genotype ($F_{2,43} = 4.23$, FDR adjusted $p = 0.056$) on average licking received. For Oxtr, homozygous G/G female offspring received lower average licking per bout than homozygous A/A female rat offspring (Figure 4a). In addition, there was a main effect of Esr1 (chr1:41590586) genotype ($F_{2,43} = 9.54$, FDR adjusted $p = 0.002$) on later-life licking provisioning. Heterozygous C/T female
rat offspring provided more licking than homozygous C/C (Tukey’s post hoc p = 0.001) and T/T (Tukey’s post hoc p = 0.001) female offspring (Figure 4b). There were no main effects of DA-related SNPs (Drd1, Drd2 or Dat) on early-life maternal licking received or provisioning.

3.4 | Gene x environment interactions

We restricted our gene x environment analyses to dopamine-related SNPs (Drd1, Drd2, Dat) because we hypothesized that the indirect relationship between early-life licking received and later-life licking provisioning would be mediated by dopaminergic activity in the maternal brain. We also restricted our outcome measures to later-life licking provisioning and DA levels in the nucleus accumbens because this was the only dependent variable that significantly correlated with later-life maternal licking provisioning. We analyzed both total licking received and average licking received as independent variables.

Drd2 (rs107017253) genotype interacted with average licking received on DA levels in the nucleus accumbens (coefficient β = −0.2643 ± 0.093, t₁,₄₂ = −2.8366, FDR adjusted p = 0.0426; Figure 5). More specifically, there was a significant effect for the A/G genotype (Effect = 0.230 ± 0.071, t = 3.2363, p = 0.0024) but not the A/A genotype (Effect = −0.0344 ± 0.060, t = −0.5700, p = 0.5718). No other gene x environment interactions were observed with total licking received or other dopamine-related SNPs. In addition, there were no significant gene x environment interactions on later-life licking provisioning.

3.5 | Moderated mediation of early-life licking received on later-life licking provisioning

Based on the reported results, we conducted a moderated mediation analysis in PROCESS (Model 7) using average licking received as the
independent variable, percent licking provisioning as the outcome-dependent variable, DA levels in the nucleus accumbens as a mediator, and Drd2 (rs107017253) genotype as a moderator. Figure 6 displays the moderated mediation statistical model with all output coefficient $\beta$ values. These $\beta$ values also reflected previous analyses with a statistically significant Drd2 (rs107017253) genotype by average licking received interaction on DA levels in the nucleus accumbens as well as a statistically significant positive correlation between DA levels in the nucleus accumbens and maternal care provisioning.

We found an indirect moderation of Drd2 genotype between average licking received and later-life licking provisioning by DA levels in the nucleus accumbens (index of moderated mediation $= -0.2301$, bootstrap 95% CI [$-0.4697$, $-0.0193$]). Specifically, there was a significant indirect effect for the A/G genotype (effect $= 0.2002$, bootstrap 95% CI [0.0185, 0.4063]) but not the A/A genotype (effect $= -0.0299$, bootstrap 95% CI [$-0.1533$, 0.0910]). This suggests that higher early-life average licking received was associated with higher DA levels in the nucleus accumbens of the maternal brain but only in female rat offspring with the A/G Drd2 genotype. Higher DA levels in the nucleus accumbens of the maternal brain of the female rat offspring with the A/G Drd2 genotype were then associated with higher later-life licking provisioning.
In this study, we investigated the role of genotype in the transmission of interindividual maternal care across generations of female rat offspring. To our knowledge, this is the first study that has examined the relationship between interindividual maternal care received and maternal care provisioning to the next generation of offspring and the genetic factors that could influence this relationship. While we did not find a direct relationship between early-life licking received and later-life licking provisioning, this relationship was indirectly affected by DA levels in the nucleus accumbens and dependent on Drd2 genotype. More specifically, female rat offspring with the A/G genotype showed a positive relationship between average licking received and dopamine levels in the nucleus accumbens of the maternal brain; there was no relationship with female rat offspring with the A/A genotype. The higher DA levels in the nucleus accumbens corresponded with higher maternal licking provisioning from postnatal days 2–9. The updated moderated mediation model is displayed in Figure 7. In addition, estrogen- and oxytocin-related SNPs affected both early-life licking received and later-life licking provisioning, suggesting that genotype can interact with the early-life maternal environment and influence later-life maternal care provisioning. Given that the early-life licking received we observed represented a continuous range of maternal care, these results suggest that genotype has an important role in the provisioning of maternal care to the next generation in the average rat mother.

We first investigated the direct relationship between early-life licking received and later-life licking provisioning and did not find any correlation between the two measures. Previous work in rats has shown that inducing early-life adversity and, in studies observing natural variations in maternal care, the outliers of maternal care received (± one standard deviation) can transmit across generations of offspring (Champagne et al., 2003; Francis et al., 1999; Gonzalez et al., 2001), demonstrating that early-life environmental influences are sufficient to alter maternal care to the next generation. However, there is substantial within-litter variation in the intergenerational transmission of maternal care, especially with female offspring born to mothers that provide average levels of licking/grooming (Champagne et al., 2003). Since we did not select for the tail ends of the normal distribution in interindividual licking received across all litters for the maternal care provisioning observations, our findings may reflect the substantial variation in intergenerational maternal care transmission within the normal distribution of maternal care received. However, other reasonable explanations for the lack of direct
effect of early-life licking received on later-life licking provisioning were that we used different observation methods (continuous coding after a brief pup separation and instantaneous sampling of undisturbed litters, respectively) and only the F₂ litters included male pups. It is possible that there would be a direct relationship between early-life licking received and later-life licking provisioning if the limitations of this study were addressed in future work.

We found that the DOPAC/DA ratio (a measure of dopaminergic activity) in the nucleus accumbens was negatively associated with later-life licking provisioning, mainly due to the positive correlation with DA levels than a negative correlation with DOPAC levels. Previous studies have shown links between maternal care provisioning and dopaminergic activity in the nucleus accumbens, especially the nucleus accumbens shell (Afonso et al., 2008, 2011; Champagne et al., 2004; Lee et al., 1999). In addition, artificially reared female rat pups show later-life elevated basal DA levels and lower pup-evoked DA levels in the nucleus accumbens and a reduction in maternal care (Afonso et al., 2011; Gonzalez et al., 2001) than normal maternally reared offspring. Additional studies are needed to explore the potential role of the nucleus accumbens and its subregions in the transmission of the effects of maternal care received at the early-life pup stage.

We detected 792,253 genetic variants that appear to segregate in the Long-Evans population, overlapping 20,508 Ensembl gene annotations. This number is likely an underestimation due to RNA-seq data being enriched in reads from transcribed regions. Of sixteen variants of interest that we verified using massarray, five were not polymorphic. The two most likely reasons for this are that either (a) the variant did not exist in our population of Long-Evans, either due to genetic drift, different suppliers, or simple change; or (b) they were false-positive calls, due to the greater difficulty in calling variants from RNA-seq data than from DNA-seq data. Notably, our Long-Evans population derived from only six breeding F₀ males, which may have lowered the genetic diversity of the F₂ offspring. However, the majority of our calls variants were confirmed (69%), and five are novel variants (83% of six novel variants probed). These findings show that this method is useful in understudied populations, which is important because a large amount of rat genetic variation is still undocumented. The data we have generated and publically shared on the genetic variation in the Long-Evans rat strain stand to be a useful resource for other researchers studying the effects of natural genetic variation and epigenetic mechanisms on phenotype.

We found that single nucleotide polymorphisms (SNPs) in candidate genes related to maternal care (oxytocin receptor and estrogen receptor alpha) affected early-life licking received or later-life licking provisioning. Previous work suggests that genetic variants in Oxt can affect infant attachment in humans (Chen et al., 2011) and later-life social attachment in prairie voles with brain region-specific effects on Oxt expression (King et al., 2016). In addition, preliminary studies have found that genetic variants in the Esr1 gene were associated with harsh parenting (Lahey et al., 2012) and could play a role in postpartum depression (Pinsonneault et al., 2013) in human populations. However, the effect of these individual SNPs on early-life licking received and later-life licking provisioning was minimal after accounting for multiple comparisons. In addition, we did not genotype the F₀ parent generation, and therefore, we were unable to account for the potential effects of the F₀ mothers’ genotype on early-life licking received. Previous work in mice has demonstrated that the genotype of the offspring can affect the amount of care they receive, termed offspring genetic effects (Ashbrook et al., 2015, 2017), which implies that the genetic profile of an organism can also affect its early-life experience and the organism is not simply a passive recipient of environmental exposures. Additional work is needed to determine biological consequences of these SNPs, and it is likely that there are additive effects of different genotypes on maternal care received and provisioning that we were unable to elucidate in this study. The role of genotype in the male lineage may also be an important consideration when studying intergenerational transmission of maternal care, as the fathers provide half their genetic profile to the offspring and play an essential role in genomic imprinting (Wilkinson et al., 2007).

We found a gene × environment interaction with average licking received and Drd2 genotype (rs107017253) on DA levels in the nucleus accumbens. We previously reported similar interactions between average licking received and the same Drd2 SNP in other DA-related behaviors (Lauby et al., 2019), but involving the A/A genotype, not A/G. It is possible that the nature of this interaction changes when the brain undergoes significant changes in the dopaminergic system, such as when a female rat is pregnant and preparing for postnatal care of her offspring (Akbari et al., 2013; Byrnes et al., 2001). Identifying other biological mechanisms that underlie these gene × environment interactions, such as linkage disequilibrium and gene expression changes, could help elucidate this discrepancy especially since the female offspring with the A/G genotype are the minority population and therefore may be statistically underpowered in observational studies. It would be important to investigate brain region-specific changes in gene expression as differences in dopamine levels in the nucleus accumbens could derive from altered function of other brain regions in the mesolimbic dopaminergic system (e.g., ventral tegmental area). Studies that investigate epigenetic mechanisms (e.g., DNA methylation) that would affect the expression of these genes would also be an important component in elucidating mechanisms underlying gene × environment interactions, as both maternal care received and the neural changes that prime mothers to provide care for offspring involve alterations in epigenetic mechanisms and gene expression (Beery et al., 2016; Cameron et al., 2008; Champagne, 2008; Stolzenberg et al., 2014).

Studies of gene × environment interactions on maternal behavior to date have focused on human cohorts. These studies have identified SNPs in oxytocin- and dopamine-related genes that interact with early-life experiences on different components of mothering behavior (Fleming & Kraemer, 2019; Jonas et al., 2013; Mileva-Seitz et al., 2012, 2013; Tombeau Cost et al., 2017) and that can be mediated by executive function (Tombeau Cost et al., 2017) or depressive
symptoms (Jonas et al., 2013). A strength of our study is application of statistical models typically used in human cohorts to rat populations, where causal biological mechanisms can be more readily examined (Jenkins et al., 2016). More specifically, we can look directly into the rat maternal brain for specific neurotransmitters, neuropeptides, and expression of relevant genes that would not be possible with human populations. This cross-species approach is important to refine hypotheses on factors important for maternal behavior that are evolutionarily conserved and has been essential to understand the maternal brain (Feldman, 2016).

Overall, this study suggests that offspring genotype for the dopamine receptor 2 gene may be an influential factor when assessing the link between maternal care received and maternal care provisioning. More broadly, offspring genotype in DA-related genes may be a critical factor involved with the development of the dopaminergic system (Lauby et al., 2019; Lovic et al., 2013). While it has been shown in previous studies that early-life maternal licking received substantially alters the functioning of the dopaminergic system, the underlying biological mechanisms are not as well-understood as other neuroendocrine systems (Stolzenberg & Champagne, 2016). Elucidating the functional consequences of the Drd2 SNP analyzed in this study will be important to understand the underlying biological mechanisms in the developmental programming of maternal care and its transmission across generations of female offspring.

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CONFLICT OF INTEREST
The authors of the manuscript have no conflicts of interest to declare.

AUTHOR CONTRIBUTIONS
SCL, PP, ASF, and POM designed the study. SCL and PP conducted the maternal care received and maternal care provisioning observations. DGA ran the variant calling analysis from Long-Evans RNA-seq datasets and created the track for the UCSC genome browser. HRM did the brain cryosectioning and liver DNA extractions for genotyping. DC conducted the HPLC. SCL and HRM analyzed the data. ASF and POM supervised the research. SCL, DGA, HRM, and POM wrote the manuscript. All authors contributed to this manuscript and approved the final version of this manuscript.

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DATA AVAILABILITY
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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Additional supporting information may be found online in the Supporting Information section.

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