INTRODUCTION

Parental care evolves to buffer offspring against environmental stress and once evolved, can be instrumental for ensuring success across a range of ecological conditions even beyond those that promoted its evolution (Clutton-Brock, 1991; Royle et al., 2012; Wilson, 1975).

In the face of rapidly changing environments, phenotypic plasticity in family interactions could contribute to species persistence by...
allowing parents to invest more when conditions are poor and for offspring to compensate when conditions exceed those that parents can or are willing to buffer. In some cases, this phenotypic plasticity may be immediately detectable as a change in behaviour. For example, many birds compensate for hot breeding season conditions by extending their time and effort at the nest, sometimes risking dehydration; some species may be immediately detectable as a change in behaviour. For example, many birds compensate for hot breeding season conditions by extending their time and effort at the nest, sometimes risking dehydration. Plasticity may also manifest in physiological responses; for instance, offspring have been shown to upregulate metabolic and immune pathways in response to the removal of a caring parent (Körner et al., 2020; Mashoodh et al., 2021; Ziadie et al., 2019) or exposure to pathogens (Körner et al., 2020). When parents and offspring are pushed to their physiological limits, altering behaviour may not be possible and maintaining phenotype stability (i.e., continuing to provide care and develop at the same level) probably depends on some combination of parental and offspring plasticity at other levels.

Transcriptomic approaches are useful for examining gene expression plasticity beneath the surface of phenotypes because gene expression is inherently plastic, and gene expression profiles can undergo profound changes while maintaining stable higher-level phenotypes (Eberwine & Kim, 2015; Nijhout et al., 2019; Rivera et al., 2021); that is, many “genetic programmes” can give rise to identical phenotypes. While most research in this area has focused on the role of plastic gene expression in maintaining physiological performance and stabilizing morphological traits under stress (Cheviron et al., 2008; Cheviron & Brumfield, 2011; Oleksiak et al., 2005; Peck et al., 2015), compensatory molecular mechanisms are also expected to underpin stability of complex behaviours across fluctuating environments (Fischer et al., 2021). This perspective is valuable to the study of parental care because parenting evolves to minimize environmental fluctuations and involves transformational changes of gene expression (Parker et al., 2015; Ray et al., 2016; Bukhari et al., 2019; Fischer et al., 2019; Lopes & de Brujn, 2021), and the ultimate configuration of expressed genes associated with care may differ depending on environment. Because parental care is inherently a social exchange and phenotypic plasticity can stem from donors and/or recipients of care, a comprehensive test of these predictions should profile the gene expression of parents and offspring simultaneously (Butler & Maruska, 2021).

Here, we performed RNA-seq on a subsocial burying beetle, *Nicrophorus orbicollis*, to explore if plasticity of gene expression helps buffer families against environmental stressors. There is a strong basis for studying the genetics of parenting and development of burying beetles (Benowitz, McKinney, Cunningham et al., 2017; Cunningham et al., 2019; Jacobs et al., 2016; Palmer et al., 2016; Parker et al., 2015; Won et al., 2018), yet the genes responsible for mounting responses to major environmental stressors while beetles are parenting/being parented remain uncharacterized. Such mechanisms are particularly relevant in the context of rapid environmental change, as they could provide targets of selection in populations unable to behaviourally avoid stressful conditions. Even less understood is the extent to which application of a secondary stressor may modify core parental responses; that is, how plasticity in behavioural gene expression buffers higher level phenotypes.

High temperatures (≥23°C) as naturally occur during the peak summer breeding months in the southeast USA. (Trumbo, 1990; Ulyshen & Hanula, 2004) impose steep reproductive and survival costs for *N. orbicollis* (Feldman, 2020; Moss & Moore, 2021; Ong, 2019; Quinby, 2016). Despite this, we found that females and males do not alter their parental care behaviour, type, or intensity when parenting at 24°C compared to at a more benign temperature, 20°C (Moss & Moore, 2021). Whether behavioural stability of parents results from gene expression changes that alter or supplement the genetic programme for parental care, or whether gene expression changes in offspring may independently compensate for thermal stress, is not known. Here, we address these possibilities while controlling for any confounding effects of differences in care. We examined the effects of two thermal environments to characterize “thermal response” genes (20°C or 24°C, the same as that investigated in Moss & Moore, 2021) and parent-offspring interactions to characterize “parenting/being parented” genes (before or during active parenting), comparing responses of mothers, fathers, and offspring to understand the similarity of each’s response. Our prediction was that thermal stressors and parental care will each elicit independent gene expression responses and that these will differ between family members (represented by the main effects of the statistical model). Further, we predicted that some genes would show distinct patterns of expression change in response to the combination of stressors to stabilize the behaviours across environments (“buffering genes” represented by the interaction term of the statistical model); specifically, higher temperatures should significantly modify expression of core genes for parenting/being parented or elicit distinct changes in gene expression during family interactions if genetic plasticity facilitates behavioural stability across environments.

2  MATERIALS AND METHODS

2.1  Study system

*Nicrophorus orbicollis* is a biparental carrion beetle that breeds on small vertebrate carcasses in woodlands ranging from southern Canada to northern Texas. At the southeastern edge of this distribution, daily mean temperatures as low as 18°C and as high as 25°C may arise over the course of the long summer breeding season (Moss & Moore, 2021). Temperatures at the upper end of this range (23–25°C) are highly challenging for parents and developing offspring, with families forced to breed under these conditions suffering increased mortality in adults and larvae and reduced clutch size larval mass (Moss & Moore, 2021). As in most members of the genus, *N. orbicollis* expresses a large repertoire of pre- and post-hatching care behaviours (Eggert & Müller, 1997; Scott, 1998). Carcass preparation begins with the removal of hair and liberal application of anal excretions, which suppress microbial growth on the prepared “brood ball”. Females oviposit throughout the soil surrounding the brood ball, and
within 2–3 days these eggs hatch and larvae migrate through the soil to the brood ball attended by their parents. At this stage, parents commence directly provisioning begging young via regurgitation. While larvae of *N. orbicollis* have relatively high starvation tolerance (mean time to starvation: ~17h), upon hatching their capacity to self-feed from a prepared carcass is extremely limited, and they rely on at least 3 h of post-hatching parental care for survival (Capodeanu-Nägler et al., 2018). This post-hatching stage is likely to be the most energetically demanding stage of care for parents as it is when attendance to offspring is highest and the stage at which we see the most sex differences of parental behaviour (Moss & Moore, 2021).

### 2.2 Study design

We used *N. orbicollis* beetles that had been collected locally and reared in the laboratory for one generation at the University of Georgia, as described in Moss and Moore (2021). Different beetles were used to generate behavioural (Moss & Moore, 2021) and transcriptomic data sets (this study). Briefly, adults were housed individually in plastic containers (9 cm diameter, 4 cm deep; Eco Products) filled with potting soil and fed organic ground beef twice weekly ad libitum. Stock breeding took place at room temperature (20 ± 0.5°C), and resulting offspring were divided evenly between two incubators starting 3 days into pupation for a thermal acclimation period prior to breeding. These incubators were programmed to long-day light conditions (reverse light: dark 14:10) on ramping temperature cycles to simulate a diurnal range: the first fluctuating between 21 and 20°C and the second between 25 and 24°C. Because breeding takes place underground in relatively thermostable conditions, however, only the lower end of the diurnal range of each thermal treatment (20 and 24°C) was used for breeding. Virgin, non-sibling beetles (aged ≥14 days post-eclosion) were paired in plastic boxes (17.2 × 12.7 × 6.4 cm; Pioneer Plastics) filled with approximately 2 cm of moistened soil and a 40–45 g thawed mouse carcass (RodentPro) and moved to either a constant dark temperature-controlled room (20°C) or incubator (24°C) corresponding to their acclimation environment. Boxes were monitored twice daily for the presence of eggs, and after 2–3 days eggs were collected into petri dishes containing damp filter paper and stored at their respective treatment temperatures. Eggs were checked every 4 h between 08:00 AM and 17:00 PM until hatching, which occurs within 2 days after laying.

In addition to manipulating the thermal environments in which care took place, we also generated within-temperature treatment controls to disentangle the effects of “active” parenting (the stage at which parents and larvae interact socially, which we refer to as parenting or direct parental care) from a general parenting state which includes behaviour such as carcass preparation that does not require social interactions and begins before larvae are present. Thus, we sampled at two time points: (1) post-larval hatching but before larvae and parents interacted (∼16 h old larvae), and (2) after the first 24 h of direct parental care. These larvae were collected directly from the petri dish immediately when found to have hatched, with parents collected simultaneously from the carcass they were preparing. Larvae of families for parenting samples, by contrast, were transferred to brood balls attended by parents when found to have hatched. These families were left undisturbed for 24 h so that direct parenting could take place before collecting both parents and offspring (24–40 h old larvae) for analysis. Parents in both treatments had initiated indirect parental care involving preparation of a brood ball for larval feeding. Timing of sampling parents before interactions coincided with the circadian switch to active parenting (Oldekop et al., 2007).

### 2.3 Sample collection, preparation, and RNA sequencing

Our design yielded 10 families at 20°C (five before parenting, five during parenting) and nine families at 24°C (four before parenting, five during parenting). We collected the heads of adults, consisting of brain and associated “fat body” tissue, as in previous studies (Parker et al., 2015). This allowed us to capture gene expression in the two tissue types that influence behaviour in insects. Because of the size of larvae at hatching and because we had no a priori predictions for the specific tissues affected by interactions with adults, we collected whole bodies. Larval samples therefore consisted of seven to 10 larvae for families before parenting and two to four larvae for families during parenting. Individual adult samples, both before and during parenting, were age-matched. Samples were flash frozen in liquid nitrogen and stored at −80°C until RNA extraction. Extractions were performed following the Qiagen RNeasy Lipid kit protocols (Qiagen). Sequencing libraries were prepared from 1.3 μg total RNA using the Illumina TruSeq mRNA Stranded Library Kit (Illumina, Inc.) according to the manufacturer’s protocol with standard Illumina adapters and primers. Sequencing was carried out on a NovaSeq 6000 platform with a 150 bp paired-end protocol, targeting 40M read pairs per sample.

### 2.4 Read mapping, transcriptome assembly, and annotation

Reads were trimmed using Trimmomatic (version 0.39; Bolger et al., 2014) to remove adapter sequences with default parameters, except low-quality bases (Phred < 15 at the leading and trailing ends and from 4-base sliding windows), and short reads (<36 bp). Reads were corrected using RCorrector with default parameters (version 1.0.3.1; Song & Florea, 2015). Quality was assessed using FastQC (version 0.11.9; http://www.bioinformatics.babraham.ac.uk/projects/fastqc/; Table S1). We first mapped our RNA-seq reads against the genome of *N. orbicollis* (Benowitiz, McKinney, Cunningham et al., 2017; Benowitiz, McKinney, Roy-Zokan et al., 2017; NCBI BioProject PRJNA371654; Methods S1). We used HISAT2 to perform the read mapping with default parameters (version 2.1.0; Kim et al., 2015). We then used StringTie to assemble the mapped RNA-seq reads into gene and isoform models (version 2.1.1; Pertea et al., 2015).
et al., 2015). We removed redundant isoforms CD-HIT (version 4.8.1; Li & Godzik, 2006) with a sequence similarity cutoff value of 95%. Finally, we assessed the completeness of the assembly using BUSCO (Insecta data set; version 5.0.0; Simão et al., 2015).

Annotation of contigs was performed using a combination of approaches. Briefly, the longest isoform of each gene was screened against the annotated and published genome of the closely related burying beetle N. vespilloides (NCBI assembly accession no. GCF_001412225.1; Cunningham et al., 2015) using Magic-BLAST (version 1.5.0; Boratyn et al., 2019). Transcripts were also fed through the "dammit" pipeline (http://db-lab.github.io/dammit/), which identifies candidate coding regions in transcripts with Transdecoder (version 2.0.1; https://hpc Illri.cgiar.org/transdecoder-software) and annotates them, drawing from multiple databases. We used the N. vespilloides proteome (Cunningham et al., 2015) as a user database to search against and also retained any hits to known protein domains in Pfam-A (version 32.0; El-Gebali et al., 2019) and orthologues in OrthoDB (version 10; Kriventseva et al., 2019). Orthologous genes were retrieved from the OrthoDB database by searching cluster IDs at the level Polyphaga. Gene ontology (GO) terms for annotated genes were assigned by using Pfam accession numbers and GO terms of OrthoDB assignment. GO terms were also supplemented using eggnOG-mapper (version 5.0; Huerta-Cepas et al., 2019) and OrthoDB searches at the Order (Insecta) level. Unique GO terms from any source were retained for a gene.

2.5 | Transcript quantification and filtering

We estimated transcript abundances for each sample separately using StringTie. A transcript count matrix was extracted with the prepDE.py script provided with StringTie and imported into R (version 4.0.3; R Core Development Team, 2019) along with sample library sizes (calculated using Picardtools, version 2.21.6; http://picard.sourceforge.net) and transcript lengths. We computed the fragments per kilobase per million (FPKM) matrix with the package (version 1.0; Alhendi, 2019). For each family member and social state (before or during parenting), we retained transcripts that were expressed >2 FPKM in more than half of samples and then combined all unique transcripts across groups in downstream analyses (Bloch et al., 2018). Finally, we used IsoformSwitchAnalyzeR (version 1.12.0; Vitting-Seerup & Sandelin, 2019) to convert transcript-level counts into gene-level counts.

2.6 | Differential expression analysis

We performed differential expression analysis using DESeq2 (Bioconductor version 1.30.0; Love et al., 2014). To cluster samples, we first applied variance stabilizing transformation using DESeq2 and performed a principal component analysis (PCA) using R’s built-in “prcomp” function. Visualization of the clustering was done using DESeq2’s “pcaplot” function and extracted principal components were regressed on each grouping factor (family member, temperature, parenting, and the statistical interactions among these factors) via analysis of variance (ANOVA using R). Significant differences among family members in the full model ANOVA led us to analyze males, females, and larvae separately. For each family member, differential expression was estimated using parametric dispersion. We used the likelihood ratio test (LRT) implemented in DESeq2 to compare the goodness-of-fit of several models for each family member to estimate differential gene expression. Two tests were used to capture gene expression changes in response to main effects. The first compared a full model fitted with both temperature and parenting (temperature+parenting) to a reduced model fitted only with parenting to identify differentially expressed genes mutually affected by temperature across parenting states (henceforth, “thermal response” genes). The second compared the same full model to a reduced model fitted only with temperature to produce a list of differentially expressed genes mutually affected by parenting across thermal environments (henceforth, “parenting” genes). The p-values that resulted from these tests were adjusted for multiple testing using the Benjamini-Hochberg correction (Benjamini & Hochberg, 1995) and genes with adjusted p < .05 were considered statistically significantly differentially expressed genes.

To assess similarity of gene expression responses to the main effects of temperature and parenting across family members, we extracted lists of overlapping genes from the intersect of male, female, and larval parenting/being parented and thermal response gene sets. To determine whether overlap was significantly greater than expected by chance given variable input sizes, we further carried out randomization tests. Briefly, 10,000 gene sets were randomly generated for males, females, and larvae by sampling from the full list of expressed genes in that family member a number equivalent to the size of the gene set being simulated. These random gene sets were used to calculate a null distribution of overlap values, and significance was calculated as the proportion exceeding the observed overlap for each pairwise comparison. While overlap of the most highly significant genes between two sets lends itself to intuitive interpretation, use of fixed thresholds can be overly stringent and lead to false negatives. Therefore, we implemented a complementary threshold-free approach: rank-rank hypergeometric overlap (RRHO) analysis. This analysis ranks DE genes according to their significance level along with the magnitude and direction of change to make qualitative assessments of concordance between complete gene sets independent of p-value cutoff. Results were visualized using the stratified method of the R package RRHO2 (version 1.0; Cahill et al., 2018), with genes ranked according to the recommended metric of −log10(p-value) * sign(log2 fold change). Finally, we assessed whether genetic programmes for parenting/being parented were altered or supplemented in response to thermal stress in two ways. First, we compared differential expression of parenting/being parented genes between the two thermal environments. Even if changes of expression of these genes are similar enough across thermal environments to be captured as main effects, the magnitude of the response may be enhanced or dampened due to thermal stress. To test this possibility, we estimated log2fold
changes of expression of parenting/being parented genes separately for each temperature treatment using the "ashr" shrinkage estimator with false discovery rate correction (Stephens, 2017) and fit a major axis regression to these data using the R package (S)MATR (Warton et al., 2006). A slope significantly different from one allows rejection of the null hypothesis that responses are generally similar in magnitude across thermal treatments and provides a more holistic comparison of the treatments. Second, we performed a differential expression analysis using the LRT approach to identify genes showing distinct responses to parenting/being parented (i.e., substantial differences in magnitude or different signs) in the presence of thermal stress (henceforth, “buffering” genes). To obtain this list, we compared a full model fitted with a statistical interaction term (temperature x parenting) to a reduced model fitted only with the main effects (temperature + parenting). To further characterize patterns of differential gene expression across social and temperature groups, we performed an ANOVA followed by post-hoc contrasts (Tukey HSD) at the level of each gene in the buffering set, comparing the magnitude and sign of differential expression at 24°C versus 20°C.

2.7 | Functional enrichment analysis

We queried the differentially expressed gene sets and the overlapping gene sets for enriched GO terms using topGO (version 2.42.0: Alexa & Rahnenfuhrer, 2020) searching for enrichment in all three GO categories: biological process, molecular function, and cellular component. The nodeSize parameter was set to 10 to remove GO terms with fewer than 10 annotated genes and only terms with >1 significant gene were retained. Significance was estimated using a Fisher’s exact test following the programme’s suggested protocol. To examine the directionality of change of enriched pathways, Z-scores were calculated following the formula of Walter et al. (2015): number of downregulated genes annotated for the term subtracted from the number of upregulated genes annotated for the term divided by the square-root of the total number of genes annotated for the term.

3 | RESULTS

Our final transcriptome assembly contained 91% of single-copy conserved nucleotide orthologues (complete: 90.5% [single copy: 76.4%, duplicated: 14.1%], fragmented: 4.2%, missing: 5.3%) and a total of 12,406 genes.

3.1 | Thermal and social conditions shape gene expression of family members

We visualized differences in gene expression among male parents, female parents, and offspring using PCA. Parents and offspring separated strongly along two primary axes (Figure 1a), which together explained 77.8% of total variance in gene expression across all samples (PC1: 73.08%; PC2: 4.76%; Table 1). Family member (parent vs. larvae) clearly separated along PC1, while separation along PC2 was driven primarily by parenting (Table 1). The interactive effect of family member and parenting and family member and temperature were significantly associated with PC2 (Table 1). Visualizing family members separately with their own dispersions further clarified how global gene expression differed among groups. Female gene expression displayed considerable overlap between thermal environments but was most divided with respect to parenting, particularly in the 20°C environment (Figure 1b). Male gene expression differentiated most strongly with temperature (Figure 1c). In larvae, behavioural groups formed distinct clusters, with parenting accounting for far greater among-group differences than temperature (Figure 1d).

3.2 | Plasticity of gene expression underpins thermal response, parenting/being parented, and buffering

We used differential expression analysis to characterize plastic changes in gene expression associated with the main effects of temperature and parenting, as well as their interaction. We identified genes showing both shared (overlapping) and distinct responses to main effects across family members, consistent with variation in physiology and behaviour between sexes and life stages. There was also gene expression variation associated with buffering of behavioural responses, where family members expressing the same behaviours in different environments showed different patterns of gene expression. These included subtle expression variation of parenting/being parented genes across temperature treatments, as well as some changes that occurred specifically in response to the interaction of temperature and parenting.

3.2.1 | Thermal response genes

Temperature induced a stronger gene expression response in males ($n = 418$ differentially expressed genes; Table S2) and larvae ($n = 487$ differentially expressed genes; Table S3) than in females ($n = 284$ differentially expressed genes; Table S4). However, male, female, and larval responses to temperature showed statistically significant overlap in terms of gene identity (male–female: $p < .001$; female–larval: $p < .001$; male–larval: $p = .006$; Figure 2a). Global concordance was pronounced between males and females (Figure 2b) – both in genes that were upregulated and downregulated in response to high temperature – whereas this signal was weak when comparing females to larvae (Figure 2c) and males to larvae (Figure 2d).

3.2.2 | Parenting genes

With the parenting main effect, the highest number of differentially expressed genes was associated with being parented, with
7189 genes differentiating larvae before and during interactions with parents (Table S5). Adults had fewer differentially expressed genes than larvae, and within adults there were sex differences. Fewer differentially expressed genes were detected in males after the arrival of larvae (n = 63; Table S6) than in females (n = 189; Table S7). Despite this, females and males showed statistically significant overlap of gene identities while parenting (Figure 2e; p = .044). When considering all expressed genes, concordance was striking but directional, with overlap between the sexes concentrated in genes downregulated in both during larval interaction (Figure 2f). Females interacting with larvae and larvae interacting with parents shared more differentially expressed genes in common (n = 115) than parenting males and parented larvae (n = 28), although this did not reach statistical significance (p = .059) and neither comparison showed global signatures of concordance of gene expression (Figure 2g,h).

3.2.3 Buffering genes

To evaluate the role of gene expression plasticity in facilitating parental buffering and offspring compensation under stressful environments, we first compared gene expression responses to parenting/being parented across temperature treatments. Among the core genes for parenting/being parented (genes that were differentially expressed between parenting states across temperatures), differential expression was significantly weaker at 24°C relative to 20°C in females (slope = 0.733; 95% CI: 0.607–0.875; p = .0007; Figure 3a,b), and did not differ significantly different between thermal treatments in males (slope = 2.088; 95% CI: 0.942–12.767; p = .067). By contrast, the slope of this regression in larvae was significantly greater than one (slope = 1.134; 95% CI: 1.121–1.148; p < .0001), suggesting that responses of larvae were somewhat stronger in magnitude at 24°C than at 20°C.
We next performed a transcriptome-wide survey for genes showing distinct responses to parenting/being parented at 24°C versus 20°C, or a significant interaction between temperature and parenting. More genes met this criterion and were differentially expressed in females ($n=79$; Table S8) than in males ($n=4$; Table S9) or larvae ($n=26$; Table S10). To test whether genes showing plasticity indeed showed stronger and broader expression changes at the more stressful temperature, the 79 buffering genes in females were subjected to closer examination (Table S11). At the benign temperature (20°C), most genes ($n=57$) showed increases of expression under active parenting, whereas at the higher temperature (24°C) only five genes showed any change in expression. Hence, most buffering genes fell into two categories: gene expression levels before parenting were equivalent between thermal treatments ($n=35$) or gene expression levels started out significantly higher at 24°C than at 20°C ($n=32$; e.g., Apolipophorin-III; Figure 3c). Because plasticity

| Variable                        | PC1 (73.08% variance) | PC2 (4.76% variance) |
|---------------------------------|-----------------------|-----------------------|
|                                 | $F_{2,55}$  | $p$-value | $F_{2,55}$  | $p$-value |
| Family member                   | 5346.224   | <.0001    | 0.624      | .540      |
| Temp                            | 4.047      | .050      | 7.073      | .011      |
| Parenting                       | 27.238     | <.0001    | 88.311     | <.0001    |
| Family member $\times$ temperature | 0.499 | .611      | 11.406     | <.0001    |
| Family member $\times$ parenting | 4.852   | .012      | 92.516     | <.0001    |
| Temperature $\times$ parenting  | 2.139      | .151      | 0.925      | .341      |
| Family member $\times$ temperature $\times$ parenting | 1.653 | .203      | 1.287      | .286      |

The Significance of Bold values indicates Significance level at 0.05

**TABLE 1** ANOVA of principal components of overall gene expression

**FIGURE 2** Concordance in differential gene expression between family members as estimated by direct overlap of differentially expressed genes (a, e) and rank-rank hypergeometric overlap (RRHO; b–d and f–h). Numbers of shared differentially expressed genes associated with (a) thermal response and (e) parenting are shown in Venn diagrams for females (red), males (blue), and larvae (yellow). Panels to the right of the Venn diagrams show the results of the RRHO analysis of all 12,406 genes for each of the respective contrasts (20°C vs. 24°C: b–d; and with vs. without parenting: f–h). The x (family member 1) and y (family member 2) axes of each plot correspond to the significance of differential expression (ranked $-\log_{10}[p$-values]) multiplied by the signed log2fold difference in gene expression for each family member-specific analysis. White boundaries demarcate the switch from down- to upregulation between behavioural groups for each family member. Pixel colour corresponds to the value of the $-\log_{10}(p$-value) from the differential expression analysis, such that hot spots in the plot designate the strength and directionality of concordance in gene expression between family members.
was temperature-dependent, gene expression levels during parenting were either significantly lower at 24°C (n = 29) or did not differ between thermal environments (n = 45), and only five genes showed significantly higher expression levels during parenting in the warmer treatment.

3.3 | Functional enrichment analysis

Functional enrichment of GO terms in the within-family member temperature contrasts revealed both shared and distinct enriched functional characteristics across females, males, and larvae. The most significantly enriched GO terms in parents were related to chitin and carbohydrate metabolism (Tables S12 and S13), whereas larvae showed the most enrichment for cellular housekeeping functions (e.g., ribosome biogenesis and rRNA processing; Table S14). Compared to the 20°C group, females, males, and larvae in the 24°C group predominantly showed downregulation of functional pathways.

Actively parenting females showed significant positive enrichment in a range of metabolic, catabolic, and biosynthetic functions compared to their respective noninteracting controls (Table S15). Forty-five of the 64 GO terms that were enriched in the female parenting gene set were also significantly enriched in the overlap between female parenting genes and larval parented genes, particularly terms related to the metabolism of organic, carboxylic, and fatty acids (Table S16). Conversely, analysis of GO terms in males revealed significant downregulation of nonoverlapping GO terms (e.g., larval cuticle patterning; Table S17). Larval functional pathways were predominantly upregulated in response to interactions with parents and spanned many categories, from cellular organization and differentiation to whole organism development (Table S18).

Finally, we examined GO term enrichment under the interaction of parenting and thermal environment. Buffering genes of
DISCUSSION

Parenting is expected to evolve to buffer environmental stress, but parenting involves an interaction between the parent and the offspring. For parenting to be effective, the family interaction needs to be predictable and the mechanisms responsible for stabilizing these interactions across variable environments are not well understood. Plasticity of gene expression could allow families to optimize dynamics for their current environment. If true, then parenting-related responses to an ecologically relevant stressor should have a gene expression signature. We test this prediction in a subsocial insect with biparental care. We found that thermal and social conditions both shape patterns of gene expression in *N. orbicollis* females, males, and larvae, and that many responses are distinct to specific family members. Further, considering parenting and thermal stress jointly revealed plasticity of gene expression. Genes that were significantly upregulated in actively parenting females (i.e., females interacting with larvae) showed weaker expression changes at a more stressful temperature, whereas differential expression associated with being parented was stronger under thermal stress. In females 79 genes responded to parenting in an entirely temperature-dependent manner (i.e., expression patterns were best explained by the statistical association between parenting and temperature). Specifically, these genes showed significant changes in expression at the benign temperature but not at the stressful temperature. Taken together with our previous findings of limited behavioural plasticity of *N. orbicollis* parents exposed to stressful temperatures (Moss & Moore, 2021), this implies that behavioural stability under stress is associated primarily with maintenance of existing genetic programmes, rather than switching to an alternative or supplemented genetic programme for the same complex behaviour.

Parental investment and offspring development are sensitive to environmental quality as poor conditions can impose competing physiological demands between parenting and thermal response. To capture and compare the gene expression and hence the molecular pathways involved in responses to thermal stress, we sampled families from a benign (20°C) and a stressful (24°C) thermal environment across the breeding cycle. While our previous study showed that *N. orbicollis* do not modify their behaviour in response to temperature (Moss & Moore, 2021), constitutive changes of gene expression of adults and larvae exposed to high temperatures across stages of care suggest that any phenotypic stabilization over this thermal range is at least partly due to the buffering action of background physiological processes. In parents, the predominant gene expression response to heat stress was downregulation, particularly of pathways involved in cuticle formation and carbohydrate metabolism (Tables S12 and S13). Similarly, larvae showed downregulation of many pathways involved in cellular housekeeping functions (Table S14). These findings are consistent with several long-term acclimation experiments reporting genome-wide downregulation in response to extreme heat stress, which could either point to an inability to mount systemic responses (i.e., due to insufficient evolutionary history of such extremes; Levine et al., 2011; Becker et al., 2018) or an adaptive molecular mechanism for metabolic compensation (Yampolsky et al., 2014). The high temperature applied here was at the upper limit but still within the range experienced in the habitat our beetles were collected (Moss & Moore, 2021), suggesting the latter as a more likely explanation. In sum, thermal stress appears to be mitigated through the regulation of different functional pathways in parents versus larvae but follows the same general pattern – tempering all gene expression responses at high temperatures – which could impose carry-over effects on gene expression underlying parent-offspring and offspring-parent interactions.

With our study, we provide a glimpse into these mechanisms by comparing the gene expression profiles of family members before and 24 h after offspring colonize the brood ball. Thousands of differentially expressed genes were detected in larvae spanning many functional categories, some due to differences of sampling ages dictated by experimental design. While we cannot disentangle the direct effects being parented per se on gene expression (Capodeanu-Nägler et al., 2018; Mashoodh et al., 2021) from indirect effects associated with the rapid development and growth occurring at this early stage (Won et al., 2018), our sampling design was biologically realistic given that orphaned *N. orbicollis* larvae usually succumb to starvation within a day (Capodeanu-Nägler et al., 2018). In comparison, changes of parental gene expression associated with having larvae to parent were subtle (~100–200 differentially expressed genes), in line with the hypothesis that the modification of behaviour within a state involves a much smaller subset of the genes required to transition between states (i.e., *N. vespilloides* transitioning into parenting from a nonreproductive state differentially express up to 650 genes; Parker et al., 2015, Cunningham et al., 2017, 2019; Bell et al., 2016; Benowitz, McKinney, Cunningham et al., 2017).

As in previous work on the related *N. vespilloides* (Parker et al., 2015), males and females parenting together showed strong concordance in overall gene expression patterns even though, as has been found for every study of parenting in male burying beetles, we detected fewer significantly differentially expressed genes for males. More so than males, however, actively parenting females showed intriguing patterns of overlap (Figure 2b) and concordance (Figure 2g) with parented larvae in terms of differentially expressed genes and pathways (i.e., males showed...
anticoncordance with larvae; Figure 2h). Given that mothers in biparental pairs appear to perform a disproportionate share of larval provisioning (Moss & Moore, 2021), we propose that these overlapping pathways, which include lipid and carboxylic acid metabolism, could play a role in self-feeding (i.e., from a shared resource) and/or nutrient exchange (i.e., mouth-to-mouth regurgitation). In either case, genes with shared or reciprocal functional roles between family members could serve as important targets of selection during adaptation to stressful environments, which may demand more efficient performance of caregivers, offspring, or both.

Finally, we characterized patterns of gene expression in response to parenting/being parented in combination with thermal stress. If plasticity of gene expression is important for maintaining behavioural stability of families across environments, then genes underlying parent-offspring interactions should be expressed differently or include more/different genes dependent on environment. The first way we tested this was by comparing the magnitude of differential expression of parenting/being parented genes between 24 and 20°C. We predicted that if plasticity of gene expression underlies the ability of parents to buffer environmental stress or for offspring to compensate independently, then responses would be more pronounced at the higher temperature. Support for this prediction was mixed: while genes for receiving care (i.e., in larvae) were differentially expressed more strongly at 24°C, genes for providing care (i.e., in females) showed less robust expression changes at 24°C than at 20°C (Figures 3a,b). As with global gene expression responses to thermal stress, plasticity in mothers may reflect adaptive metabolic compensation, whereas plasticity in offspring may be a correlated response to compensate for increased parental restraint.

Another possibility is that plastic responses to parenting/being parented mediated by thermal stress involve completely different genes to those captured by the main effects of parenting. If true, then many differentially expressed genes should be captured by the interaction between parenting and temperature, and changes in expression of these genes before and during parenting should be more pronounced and/or change their sign at the higher temperature. Overall few genes satisfied this condition in any family member. While not entirely surprising given that interaction terms have lower statistical power than main effects, this suggests that individuals in the same behavioural state but at a different temperature still rely on more-or-less the same genetic programme. The largest gene expression response occurred in heat stressed actively parenting mothers, and closer inspection of this “buffering” gene set (n = 79) revealed the nature of this plasticity. Genes that were differentially expressed at 20°C (i.e., significantly increased in expression during parenting) did not change expression or a temporal shift followed by sustained expression (i.e., increased before parenting but then showed no further change during parenting; Figure 3c) at 24°C. Thus, while temperature may independently influence the expression of some parenting genes, the effects are not synergistic, and rather than amplify differences between behavioural states, thermal stress appears to have a tempering effect.

Our inferences based on differential expression analysis were further corroborated by functional enrichment analysis of maternal buffering genes, which revealed significant downregulation of pathways involved in fatty acid oxidation and lipid processing (Table S18). Lipids are the principal macronutrient used for fueling energetically intensive behaviour in insects (e.g., flight: Canavoso et al., 2003), yet previous efforts to link post-hatching care (i.e., presumed to be the costliest form of care) to increased lipid metabolism in burying beetles found no relationship (Benowitz, McKinney, Roy-Zokan, et al., 2017). We suggest that females restrict the use of stored energy even more in an energetically demanding environment, implying that the combined stressors of temperature and parenting could directly mediate trade-offs between current and future reproduction. In line with this, actively parenting cichlids exposed to stressful levels of noise show pronounced gene expression changes in brain regions responsible for homeostatic functioning, suggesting that stressful environments may trigger a neural switch from offspring-promoting to self-promoting behaviours (Butler & Maruska, 2021).

Offspring of stressed mothers in our study in turn upregulated genes involved in forming the larval serum protein complex—a structure that sequesters amino acids early in development for the synthesis of adult proteins (Chrysanthis et al., 1994). Whether this gene expression response by larvae may help compensate for constrained parental investment or could simply be an indirect effect of plasticity in mothers remains to be seen. What is clear from our study is that parental gene expression responses to larvae are tempered at higher temperatures but generally involve the same genes, meaning that the adaptive function most likely fulfilled by plasticity of gene expression is metabolic compensation. Thus, while burying beetles at their southern range limit regularly encounter high breeding season temperatures and future conditions will probably become more extreme, adaptation appears to be proceeding via selection on the maintenance, rather than the replacement or supplementation of genetic programmes underlying family interactions.

**AUTHOR CONTRIBUTIONS**

Jeanette B. Moss and Allen J. Moore conceived and designed the study. Jeanette B. Moss, Elizabeth C. McKinney, Christopher B. Cunningham, and Allen J. Moore collected and analysed the data. Jeanette B. Moss, Christopher B. Cunningham, and Allen J. Moore wrote the manuscript with input from Elizabeth C. McKinney. All authors gave final approval for publication.

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CONFLICTS OF INTEREST

The authors declare no conflicts of interest.

OPEN RESEARCH BADGES

This article has earned an Open Data Badge for making publicly available the digitally-shareable data necessary to reproduce the reported results. The data is available at [provided https://doi.org/10.5061/dryad.4qrf6q4d4].

DATA AVAILABILITY STATEMENT

All sequencing data have been deposited in NCBI GenBank under the BioProject ID PRJNA715430. Raw sequence reads have been deposited in the Sequence Read Archive (SRA) under accession numbers SAMN18349415–SAMN18349472 and the assembled transcriptome has been deposited in the Transcriptome Shotgun Assembly (TSA) Sequence Database under submission SUB9312680. Supporting data and R scripts are deposited as a Data Dryad Project: https://doi.org/10.5061/dryad.4qrf6q4d4

BENEFITS GENERATED

Benefits from this research accrue from the sharing of our data and results on public databases as described above.

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**SUPPORTING INFORMATION**

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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