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The unique effects of maternal and paternal depressive symptoms on youth’s symptomatology:

Moderation by family ethnicity, family structure, and child gender.

*Development and Psychopathology.*
Abstract

Drawing on five waves of longitudinal data from 392 families (52% female; \( M_{\text{age}_W1} = 12.89, SD = .48; M_{\text{age}_W5} = 21.95, SD = .77; \) 199 European American and 193 Mexican American families; 217 intact and 175 stepfather families), this study documented transactional relations of mothers’ and fathers’ depressive symptoms with youth’s symptomatology from early adolescence to young adulthood. Trait and time-varying cross-lagged models revealed that both mothers’ and fathers’ between- and within-person differences in depressive symptoms were associated with youth’s internalizing and externalizing symptoms. However, whereas each parent’s depressive symptoms uniquely contributed to youth’s internalizing symptoms, only mothers’ depressive symptoms influenced youth’s externalizing symptoms. Although reciprocal effects of youth’s internalizing symptoms on parents’ depressive symptoms were not significant, youth’s externalizing symptoms predicted changes in mothers’ depressive symptoms over time. Moderation analyses revealed distinct transactional patterns by family ethnicity and child gender, but not by family structure. This study revealed dynamic transactions among family members’ symptomatology that point to opportune times and targets for intervention efforts aimed at mitigating the negative impact of parents’ depressive symptoms on youth’s adjustment.

Keywords: adolescence, ethnicity, externalizing, family structure, gender, internalizing, parental depressive symptoms, trait-invariant, time-varying
The adverse effects of parental depressive symptomatology on youth adjustment have been consistently emphasized in the literature (e.g., Connell & Goodman, 2002; Natsuaki et al., 2014). However, extant research on the effects of parental depressive symptoms features several significant limitations. First, prior studies have tended to focus on depressive symptomatology of either mothers (e.g., Hammen, Brennan, & Kennan-Miller, 2008; Natsuaki et al., 2014) or, to a lesser degree, fathers (e.g., Kane & Garber, 2004), but not both, despite evidence that parental depressive symptoms may have additive effects on child development (e.g., Brennan, Hammen, Katz, & Le Brocque, 2002; Reeb, Conger, & Wu, 2010). Relatedly, relative to the wealth of literature examining the effects of maternal depressive symptoms on child development (e.g., Corona, Lefkowitz, Sigman, & Romo, 2005; Natsuaki et al., 2014), there is a dearth of knowledge about paternal depression effects (e.g., Cummings, Keller, & Davies, 2005; Kane & Garber, 2004). This developmental imbalance is problematic for understanding parental depression effects generally, and particularly in later development as paternal influences become more salient amidst waning physical dependence on the mother and waxing paternal involvement in parenting (Connell & Goodman, 2002). Second, prior studies have examined infants and young children (e.g., Cummings et al., 2005; Low & Stocker, 2005) more often than adolescents (e.g., Elgar, Mills, McGrath, Waschbusch, & Brownridge, 2007; Reeb et al., 2010) or young adults (e.g., Hammen et al., 2008; Rohde, Lewinsohn, Klein, & Seeley, 2005). This pattern is concerning given the dramatic changes in youth’s social contexts across the transition from adolescence to young adulthood as platonic and romantic peer relationships take on increased salience amidst declining parental influences (Arnett, 2000; Furman & Buhrmester, 1992). Third,
across parents and time periods, most empirical evidence linking parental depressive symptomatology to development derives from cross-sectional designs as implemented in largely clinical and predominantly European American samples. As such, there has been limited consideration afforded to the reciprocal influence of youth’s symptomatology on parental depressive patterns over time (Gross, Shaw, Moilanen, Dishion, & Wilson, 2008) and certainly not within diverse groups. Together, these gaps in the literature have stymied efforts to understand the impact of parental depressive symptoms on youth adjustment from adolescence to adulthood and in diverse populations.

The current investigation sought to fill these knowledge gaps by evaluating concurrent and prospective relations between both mothers’ and fathers’ depressive symptoms on youth’s symptomatology from adolescence to young adulthood in a community sample of European American and Mexican American families. In general, studies of parental psychopathology, and of parental depression effects in particular, have not yet documented when and for whom hypothesized relations between parent and child symptomatology will be greatest. Therefore, this study evaluated whether the effects of mothers’ and fathers’ depressive symptoms on youth’s symptomatology differed by family ethnicity, family structure, and child gender.

Parental Depression in the Family System

This investigation drew on the principles of family systems theory to understand whether, when, and for whom mothers’ and fathers’ depressive symptoms influence youth’s symptomatology across adolescence and into young adulthood. Family systems theory holds that every individual is both influenced by, and reciprocally influences, other members in the hierarchical family structure (Cox & Paley, 1997, 2003). Moreover, drawing on the tenets of bioecological theory (Bronfenbrenner & Morris, 2007), the family is recognized as nested within
broader developmental and cultural systems. Therefore, this study examined the relative influence of early versus later parental symptom exposure, and the moderating effects of family ethnicity (i.e., European American versus Mexican American), family structure (i.e., intact versus stepfather family), and youth gender on pathways between parents’ depressive symptoms and youth’s symptomatology.

**Parental Depression Effects on Youth Symptomatology**

Research on parental depressive symptoms has documented both short- and long-term effects on child development. For example, in a rare longitudinal study, Cummings and colleagues (2014) found that both baseline levels of parental depressive symptoms and symptom growth over time were associated with higher rates of emotional and behavioral problems in early childhood and adolescence. Similarly, Elgar and colleagues (2007) found that both the initial level and change in mothers’ and fathers’ depressive symptoms (as composited to yield a global index of parental depression) across ages 10 and 15 predicted both more internalizing and externalizing symptoms and fewer prosocial behaviors in adolescence. Given these and other studies (e.g., Cummings et al., 2005; Goeke-Morey & Cummings, 2007; Reeb et al., 2015), the current investigation sought to extend prior findings drawn (primarily) from early childhood and adolescence to examine the contribution of mothers’ and fathers’ depressive symptoms to youth’s internalizing and externalizing symptomatology from adolescence to young adulthood.

Research documenting the effects of mothers’ depressive symptoms on youth adjustment is far better developed than the literature examining the effects of fathers’ depressive symptoms on children’s functioning. However, as discussed earlier, additional gaps in the literature remain with most of the research on maternal depression effects centered on young children (for exceptions, see Hammen et al., 2008; Reeb & Conger, 2009), using cross-sectional designs (for
exceptions, see Cummings et al., 2014; Elgar et al., 2007; Rohde et al., 2005) in predominantly
European American samples (for exceptions, see Corona et al., 2005; Huang, Costeines, Kaufman, & Ayala, 2014). Notwithstanding these limitations, mothers’ depressive symptomatology has been associated with a range of negative child adjustment outcomes, including elevated rates of internalizing and externalizing symptoms (Elgar et al., 2007; Goeke-Morey & Cummings, 2007), decreased academic achievement (Murray et al., 2010), and poor social competence (Cummings et al., 2005). Although less common, studies with adolescents and young adults point to similarly negative effects on youth adjustment (Hammen et al., 2008; Rohde et al., 2005).

Although fathers serve distinct and important roles in the lives of their children (Parke, 1996), research on the effects of fathers’ mental health and behaviors on child adjustment is sparse in comparison to research on mothers (e.g., Connell & Goodman, 2002; Phares & Compas, 1992). The need for increased consideration of paternal depression effects is supported by recent research (e.g., Reeb & Conger, 2009; Reeb et al., 2010) and shifting social ecologies that render fathers increasingly involved in their children’s lives generally (Cabrera, Tamis-LeMonda, Bradley, Hofferth, & Lamb, 2000), and particularly as children age (Connell & Goodman, 2002). Importantly, fathers evidence similar rates of depression during the child rearing years as their female counterparts (Wilson & Durbin, 2010), and available research on fathers’ depressive symptoms suggests that fathers’ symptomatology evidence significant effects on children’s internalizing symptoms, such as depression and anxiety (Connell & Goodman, 2002; Phares & Compas, 1992), and particularly strong impacts on children’s externalizing problems, such as ADHD, delinquency, and conduct disorder (e.g., Kane & Garber, 2004; Phares & Compas, 1992).
Overall, extant literature supports the salience of both mothers’ and fathers’ depressive symptoms for understanding youth’s internalizing and externalizing symptomatology. Studies examining separate models with mothers or fathers indicate that the magnitude of the effect between mothers’ depressive symptoms and youth’s internalizing symptoms tends to be larger than the effect between fathers’ depressive symptoms and youth’s internalizing symptoms with this imbalance persisting from early childhood to adulthood (e.g., Connell & Goodman, 2002; Natsuaki et al., 2014). In contrast, a recent meta-analysis indicated that the magnitude of mothers’ and fathers’ depression effects on children’s externalizing symptoms tend to be similar, with a few studies indicating that fathers’ depressive symptoms are more strongly related to children’s externalizing symptoms than are mothers’ depressive symptoms, especially during adolescence (Connell & Goodman, 2002).

Unfortunately, across studies of either maternal or paternal depressive symptoms, researchers have rarely considered the implications of the co-parent’s symptomatology on child adjustment. Of note, among the few studies to examine paternal depression effects, Reeb and Conger (2009; Reeb et al., 2010) found that fathers’ depressive symptoms were associated with early adolescents’ adjustment even after controlling for mothers’ depressive symptoms. The current investigation addressed the ongoing need for research on maternal and paternal depression effects on development, particularly in adolescence and young adulthood, to elucidate the specific influence of mothers’ and fathers’ depressive symptoms on youth’s internalizing and externalizing symptomatology.

Youth Symptomatology Effects on Parents’ Depressive Symptoms

Empirical work with young children has shown that parents with children who engage in externalizing behaviors are more likely to report higher levels of child-related stress and
acknowledge negative effects of children’s behavior problems on their social life and feelings toward parenting (Donenberg & Baker, 1993; Serbin, Kingdon, Ruttle, & Stack, 2015). However, relative to research on the effects of parents’ psychological functioning and caregiving behavior on children’s adjustment, fewer studies have examined the transactional effects of children’s behavior problems on parenting (e.g., Davidov, Knafo-Noam, Serbin, & Moss, 2015), and even fewer researchers have examined how children’s behavior may influence parents’ mental health (e.g., Gross et al., 2008). To address this gap, this investigation drew on repeated measures of both youth’s and parents’ symptomatology to examine if and how youth’s internalizing and externalizing symptoms affected mothers’ and fathers’ depressive symptoms across adolescence and into young adulthood. Despite the dearth of prior literature, we anticipated that youth’s internalizing and externalizing symptoms would predict elevated rates of depressive symptoms in mothers and fathers.

**Parental Depression Effects on Youth Symptomatology: Moderating Factors**

As reviewed earlier, the family system is situated within broader developmental and cultural systems that may qualify the expression and/or impact of parenting processes (Bronfenbrenner & Morris, 2007). The current study evaluated the moderating influence of three contextual factors on the predicted relations between parents’ depressive symptoms and youth’s symptomatology.

First, we evaluated the moderating influence of family ethnicity given that both parental psychopathology and parenting practices are shaped by societal and cultural belief systems (Marsiglio & Cohan, 2000). Although most studies on parental depression effects have employed predominantly European American samples (e.g., Cummings et al., 2014; Reeb et al., 2010), some research suggests that parental depressive symptoms have stronger relations with negative
parenting and, by extension, poorer youth adjustment in non-European American families (Veneziano, 2000; Wilson & Durbin, 2010). With regard to Mexican American families specifically, parental depression may have particularly strong effects on youth adjustment due to the heightened valuation of familial support, respect (i.e., *respeto*), closeness (i.e., *familism*), and obligation (Fuligni, Tseng, & Lam, 1999). Although mothers’ depressive symptoms are related to increased internalizing and externalizing symptoms among Latinx youth (Corona et al., 2005), this study was among the first to investigate these relations as compared to European American youth.

Second, we evaluated transactions between parents’ and youth’s symptomatology in both intact and stepfamilies. Prior research suggests that children in stepfamilies experience poorer developmental outcomes than children in intact families, including higher rates of internalizing and externalizing symptoms (Cherlin & Furstenberg, 1994; Perez-Brena, Cookston, Fabricius, & Saenz, 2012). However, beyond this main effect, the current study was among the first to examine if and how stepfamily structure, specifically stepfather families versus intact biological families, may influence relations between parents’ and youth’s symptomatology.

Third, we explored the moderating influence of child gender in light of mixed findings on the effect of child gender on relations between parental depressive symptoms and youth adjustment. Most studies with younger samples have shown that maternal psychopathology is more strongly related to internalizing problems in both boys and girls, whereas paternal psychopathology is more strongly related to externalizing problems, especially for girls (Connell & Goodman, 2002). Additional evidence suggests that the nature of these moderating effects may change over time, with boys being more susceptible to negative family environments during childhood, and girls being more vulnerable than boys in adolescence (Reeb & Conger, 2009).
Study Overview

The current investigation evaluated the unique contributions of mothers’ and fathers’ depressive symptoms to youth’s internalizing and externalizing symptoms, as well as the reciprocal effects of youth’s symptomatology on mothers’ and fathers’ depressive symptoms from adolescence (age 12) to young adulthood (age 22) within and across groups based on family ethnicity, family structure, and child gender. First, we hypothesized that higher levels of mothers’ and fathers’ depressive symptoms would each contribute to elevated rates of youth’s internalizing and externalizing symptoms across adolescence and young adulthood. We further predicted that mothers’ depressive symptoms would be more strongly related to youth’s internalizing symptoms, whereas both mothers’ and fathers’ depressive symptoms would be related to youth’s externalizing symptoms. Given research indicating that both baseline levels and change in parents’ depressive symptoms over time are related to youth’s adjustment (Cummings, Cheung, Koss, & Davies, 2014; Elgar et al., 2007), we hypothesized that both proximal and distal parental depressive symptoms would be uniquely related to youth’s internalizing and externalizing symptoms. Second, building on the robust body of literature supporting bi-directionality in parent-child relationships (e.g., Davidov et al., 2015), we hypothesized that youth’s symptomatology would influence mothers’ and fathers’ depressive symptoms with youth’s externalizing symptoms having a greater impact than internalizing symptoms on parents’ mental health. Third, we hypothesized that the effects of mothers’ and fathers’ depressive symptoms on youth’s symptomatology would vary by family ethnicity, family structure, and child gender. Specifically, we hypothesized that effects from parental depressive symptoms to youth’s adjustment, and from youth’s adjustment to parental depressive symptoms, would be stronger for Mexican American families in comparison to European
American families given the relatively higher degree of cohesion and interdependence among Mexican American families as compared to European American families. Although research suggests that children fare worse in stepfamilies than in intact families (Cherlin & Furstenberg, 1994; Coleman, Ganong, & Fine, 2000), we hypothesized that the link between both parents’ depressive symptoms and youth’s adjustment would be stronger in intact families given the shared genetic and environmental history in biological families. However, we did not expect that family structure would significantly influence reciprocal relations from youth’s symptomatology to parents’ depressive symptoms. Finally, we hypothesized that child gender would moderate associations between parents’ depressive symptoms and youth’s internalizing and externalizing symptoms. Some theorists have argued that mothers tend to have more influence on their adolescent daughters, whereas fathers have more influence on their sons during adolescence (Hill & Lynch, 1983). However, other evidence points to the disproportionate salience of cross-gender effects (e.g., Amato, 1994). Given mixed findings regarding the moderating influence of child gender, the present analyses were exploratory with regard to both parental depression effects on youth’s adjustment and reciprocal contributions of youth’s symptomatology to mothers’ and fathers’ depressive symptoms.

**Method**

**Participants and Procedures**

Families ($N = 392$; 52% female children) participated in a five-wave, dual-site longitudinal study that was conducted in Phoenix, Arizona and Riverside, California (see Stevenson et al., 2014 for full description of recruitment procedures). The study targeted families who were of European or Mexican descent and included an adolescent who was enrolled in 7th grade. All three participating family members were required to be from the same ethnic
background, and families were recruited to include both intact families (i.e., two biological parents in the household) and stepfather families (i.e., a biological mother and a male romantic partner who was acting as a “father figure” to the child in the residence). The father and the mother were not required to be legally married, but the household structure had to be in place for more than one-year. The resulting sample included 110 European American intact families (96.36% married), 89 European American stepfather families (75.28% married), 107 Mexican American intact families (94.39% married), and 86 Mexican American stepfather families (44.19% married).

Participants completed a battery of assessments at the research site or via phone in their preferred language (English or Spanish) across five data waves. Following a cohort sequential design, assessments began when the adolescent was enrolled in 7th grade ($M_{age,W1} = 12.89, SD = .48$) and continued through young adulthood ($M_{age,W5} = 21.95, SD = .77; N = 276$), with intervening assessments at wave 2 ($M_{age,W2} = 13.89, SD = .76; N = 365$), wave 3 ($M_{age,W3} = 15.53, SD = .65; N = 321$), and wave 4 ($M_{age,W4} = 19.68, SD = .70; N = 287$). Each family member received monetary compensation for their time. All procedures for this study were approved by the Institutional Review Boards of the participating universities.

Of the 392 families interviewed at Time 1, 79.8% ($n = 312$) were legally married, and there were significant differences in marriage rates across subgroups, $\chi^2(3) = 100.87, p < .001$, such that those in stepfather families were less likely to be legally married, especially among Mexican American stepfather families. The majority of mothers (67%; $n = 262$) and (step)fathers (67.5%; $n = 264$) were born in the United States. Mothers and (step)fathers who were born outside the United States had resided in the country for an average of 15.03 years ($SD = 8.01$) and 16.25 years ($SD = 8.11$), respectively, $t(238) = 1.17, ns$. The annual adjusted family income
ranged from $8000 to over $100,000, with a mean of $67,410.06 (SD = $47,194.79), though 19.6% of the families earned below $35,000 per year. There was no significant difference in family income between intact (M = 66,705.17, SD = 47,151.39) and stepfather families (M = 68,362.45, SD = 47,489.87), t(389) = .34, ns. However, European American families reported higher household income (M = 86,678.08, SD = 54,392.10) than Mexican American families (M = 47,514.62, SD = 26,588.13), t(289.79) = 9.09, p < .001.

Across the five data waves, 377 (96.2%) of the families completed two or more assessments. With the exception of youth’s depressive symptoms and externalizing behavior, there were no significant differences across all study variables at Wave 1 between families who completed two or more assessments and those who did not. Youth in the 15 families that did not participate in two or more interview assessments were more likely to report higher rates of depressive symptoms, t(14.563) = 2.09, p = .055, and externalizing behavior, t(14.386) = 2.28, p = .038.

Measures

**Parental Depressive Symptoms.** Mothers’ and fathers’ depressive symptoms were assessed using the Hopkins Symptom Checklist (HSC; Derogatis, Lipman, Rickels, Uhlenhuth, & Covi, 1974). Items (e.g., In the past month, how often have you had a poor appetite) were rated on a 4-point Likert type scale (1 = not at all to 4 = extremely), with higher scores indicating higher levels of depressive symptoms. With the exception of wave 2, all ten items for the depression scale were administered to both mothers and (step) fathers at each data wave. Only three items (i.e., In the past month, how often have you been feeling hopeless about the future, how often have you been feeling blue, and how often have you been feeling no interest in things) were administered at the second time point. Bivariate correlations between a composite variable
of the three items and a composite variable of all ten items at the other waves indicated the two scales were highly related for mothers ($r_s = .898$ to $.921$) and fathers ($r_s = .882$ to $.909$).

Moreover, these correlations remained strong for mothers ($r_s = .770$ to $.813$) and fathers ($r_s = .672$ to $.755$), even when the three items shared across measures were correlated with the seven unique items within each wave. Given the consistently strong relations between the short- and full-scale scores, with and without the shared items, the 3-item scale was used to assess mothers’ and fathers’ depressive symptoms across waves. The reliabilities for the three items were acceptable for mothers ($\alpha = .726$ to $.817$) and fathers ($\alpha = .667$ to $.744$).

**Youth’s Symptomatology**

**Internalizing Symptoms.** Youth’s internalizing symptoms were assessed by self-reports. At waves 1 through 3, depressive symptoms were assessed using items from the Child Depression Inventory (CDI; Kovacs, 1992). Eight items (e.g., in the past month, things bothered me) were scored on a 3-point scale (1 = *I did not feel alone* to 3 = *I feel alone all the time*), with higher scores reflecting higher levels of depressive symptoms ($\alpha = .652$ to $.718$). At waves 1 through 3, youth’s anxiety was assessed using items from the Revised Children’s Manifest Anxiety Scale (RCMAS; Reynolds & Richmond, 1979). Seven items (e.g., in the past month you worried about what was going to happen) were rated on a dichotomous scale (1 = *Yes* to 2 = *No*) and were recoded so that higher values indicated more anxiety symptoms ($\alpha = .651$ to $.688$). Both the CDI and RCMAS were abbreviated for use in this study due to time constraints.

Employing data from the full CDI and RCMAS scales administered in prior work (Wolchik et al., 2000), stepwise regression analyses were used to identify the current subsets of scale items that accounted for 90% of the variance in the full scale scores (see Schenck et al., 2009 for full description). At waves 4 and 5, youth’s internalizing symptoms were assessed using the 18 items...
(e.g., I feel lonely) from the anxious/depressed subscale of the Adult Self Report (ASR; Achenbach, 1991b). Items were rated on a 3-point scale (1 = not true to 3 = very true or often true; \( \alpha = .852 \) to .859).

**Externalizing Symptoms.** Youth’s externalizing symptoms were assessed by self-reports. At waves 1 through 3, youth reported on their externalizing symptoms using 12 items (e.g., in the past month, I destroyed things belonging to others) from a modified version of the aggression and delinquency subscales of the Behavior Problems Index (Peterson & Zill, 1986). These items were rated on a 3-point scale from 1 (not at all true) to 3 (very true; \( \alpha = .751 \) to .831). At waves 4 and 5, youth’s externalizing symptoms were assessed using the 35 items (e.g., I damage or destroy my things) from the aggressive, rule-breaking, and intrusive behavior subscales on the Adult Self Report (ASR; Achenbach, 1991b). Items were rated on a 3-point scale (1 = not true to 3 = very true or often true; \( \alpha = .890 \) to .894).

**Data Analytic Plan**

All analyses were conducted in Mplus 6.12 (Muthén & Muthén, 1998-2011) to account for missing data using Full Information Maximum Likelihood. Prior to the cross-lagged panel model analyses, longitudinal invariance models were estimated to evaluate whether the different measures of youth’s internalizing (i.e., CDI, RCMAS, ASR) or externalizing symptoms (i.e., BPI, ASR) were assessing the same constructs across time (see Author, under review). For the cross-lagged panel model analyses, factor score estimates were computed from weak factorial invariance models for internalizing and externalizing symptoms to reduce computational burden and possible overestimation. Factor scores are estimates of the underlying score on the latent variable or factor for each observation or participant (Brown, Hendrix, Hedges, & Smith, 2011).
A series of cross-lagged panel models evaluated the effects of mothers’ and fathers’ depressive symptoms on youth’s internalizing and externalizing symptoms from adolescence to young adulthood, as well as reciprocal effects from youth’s symptomatology to mothers’ and fathers’ depressive symptoms. Research has shown that failing to disaggregate between- and within-person differences in cross-lagged panel analyses can yield erroneous conclusions about causal patterns and/or biased and difficult to interpret parameters or coefficients (Berry & Willoughby, 2016; Hamaker, Kuiper, & Grasman, 2015). Therefore, each symptom model included a global trait factor and five state (time-varying) factors to disaggregate trait-invariant differences (e.g., between-person effects) from time-varying differences (e.g., within-person effects) for mothers, fathers, and youths. In each model and for all three reporters, the trait factor, and the five time-varying factors were fixed at zero. The variances of the factor score estimates were fixed at zero, whereas the variances for the trait factor and the time-varying factors were freely estimated. The loadings from the trait factor to the factor score estimates at each wave were fixed to one. The stability coefficients for the time-varying factors were also included in this model (e.g., the time-varying internalizing factor at wave 1 predicted the time-varying internalizing factor at wave 2). Covariances between nonadjacent waves were fixed at zero (e.g., the time-varying internalizing factor at wave 1 was not associated with the time-varying internalizing factor at wave 3). Separate cross-lagged panel models were estimated to evaluate reciprocal effects of mothers’ and fathers’ depressive symptoms on youth’s internalizing or externalizing symptomatology from adolescence to young adulthood.

Multigroup analyses tested for differential effects by family ethnicity, family structure, and child gender on bidirectional associations between mothers’ or fathers’ depressive symptoms and youth’s internalizing and externalizing symptoms. Sattora’s (2000) likelihood ratio chi-
square difference test evaluated comparative fit across each pair of nested models. However, given that the likelihood ratio test is influenced by large sample size (Browne & Cudeck, 1993), additional practical fit indices were examined, including the Tucker Lewis Index (TLI; Tucker & Lewis, 1973), comparative fit index (CFI; Bentler, 1990), root mean square error of approximation (RMSEA; MacCallum, Browne, & Sugawara, 1996), and standardized root mean square residual (SRMR; Hu & Bentler, 1999). Good model fit was indicated by TLI and CFI values greater than .95, and RMSEA and SRMR values below .08 (Hu & Bentler, 1999; MacCallum et al., 1996).

Results

Internalizing Symptoms

See Table 1 for descriptive information regarding mothers’ and fathers’ depressive symptoms. In all models, a stable trait latent construct (Kenny & Zautra, 2001) was created to capture the between-person differences for each reporters’ symptoms. In addition, the time-varying components for each reporter’s symptoms were fixed according to a simplex pattern (e.g., youth’s wave 1 internalizing symptoms predicted their wave 2 internalizing symptoms, but not the nonadjacent waves). At each time point, the time-varying variables for all reporters were correlated. In addition to the inclusion of stability coefficients and cross-lagged influences on each reporter, the global trait factor for each reporter was correlated with the other two reporters’ global trait factor (e.g., mothers’ trait-like depressive symptoms were correlated with youth’s trait-invariant internalizing symptoms and father’s trait-like depressive symptoms). For wave 5 youth’s internalizing symptoms, both parents’ proximal effects (e.g., wave 4) and distal effects (e.g., wave 1) were allowed to predict changes in youth’s time-varying internalizing symptoms. Changes in all models refer to time-varying variances that reflect both within-person variances as
well as wave-specific variability. Furthermore, the notations for covariances are denoted as “c” and the unidirectional path estimates are denoted as “b.”

The baseline cross-lagged model (Model A) fit the data well, $\chi^2(46) = 67.851, p = .020$, RMSEA = .035 [.014, .052], CFI = .991, TLI = .979, SRMR = .033. However, in Model B, the covariances between reporters for later waves were removed from the model to be consistent with traditional cross-lagged panel designs, $\chi^2(58) = 87.556, p = .007$, RMSEA = .036 [.019, .051], CFI = .987, TLI = .977, SRMR = .036. Model B did not differ significantly from Model A, $\Delta \chi^2(12) = 19.705, p = .073$, suggesting that the later covariances did not contribute significant information to the model.

Model B revealed both trait-invariant and time-varying differences in youth’s internalizing symptoms across time with the standardized coefficients for the global trait factor ranging from .674 to .741 (see Figure 1). These trait-like differences in youth’s internalizing symptoms appeared consistent across time. However, youth also showed significant time-varying changes in their internalizing symptoms after removing their trait-like differences. Across early and middle adolescence (i.e., waves 1-3), youth’s internalizing symptoms varied across time, $b = .459$ to $b = .217$, and this time-varying component of internalizing symptoms became very stable between late adolescence and young adulthood (i.e., waves 4-5), $b = .866$.

For mothers’ depressive symptoms, only one time-varying coefficient was significant. Specifically, mothers’ time-varying depressive symptoms at wave 4 were positively associated with their time-varying depressive symptoms at wave 5, $b = .232$, $SE = .070, p < .001$. These findings suggest that the variation in mothers’ depressive symptoms can be explained by between-person differences. A similar pattern was found for fathers’ depressive symptoms. Fathers’ time-varying depressive symptoms at wave 3 were negatively associated with fathers’
time-varying depressive symptoms at wave 4, $b = -.203$, $SE = .109$, $p = .061$. However, there were no other significant time-varying associations across waves, suggesting that most of the variation in fathers’ depressive symptoms was explained by stable between-person differences at the trait level.

Correlations between the global trait factor among the three reporters revealed that mothers’ and fathers’ trait-invariant differences in depressive symptoms were related to youth’s trait-invariant internalizing symptoms across time. Specifically, youth’s trait-invariant differences in internalizing symptoms were positively associated with mothers’ ($c = .176$, $SE = .047$, $p < .001$) and fathers’ ($c = .111$, $SE = .041$, $p = .006$) trait-invariant depressive symptoms. Similarly, mothers’ trait-invariant differences were positively associated with fathers’ trait-invariant differences in depressive symptoms, $c = .304$, $SE = .078$, $p < .001$.

At wave 1, the time-varying covariance between mothers’ and fathers’ depressive symptoms was significant, $c = .332$, $SE = .092$, $p < .001$. In contrast, youth’s time-varying internalizing symptoms did not correlate significantly with mothers’ ($c = .049$, $SE = .056$, $p = .385$) or fathers’ ($c = .048$, $SE = .047$, $p = .380$) time-varying depressive symptoms at wave 1. The cross-lagged results revealed that mothers’ ($b = .034$, $SE = .017$, $p = .046$) and fathers’ ($b = -.04$, $SE = .020$, $p = .042$) time-varying depressive symptoms at wave 1 predicted changes in youth’s time-varying internalizing symptoms at wave 5 (i.e., distal effects). However, mothers’ and fathers’ time-varying depressive symptoms at wave 4 did not predict changes in youth’s time-varying internalizing symptoms at wave 5 (i.e., proximal effects). Furthermore, there were no significant associations between both parents’ depressive symptoms and youth’s internalizing symptoms at earlier waves. Fathers’ time-varying depressive symptoms at waves 2 and 3 predicted lower levels of mothers’ time-varying depressive symptoms at wave 3, $b = -.178$, $SE = .
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.095, \( p = .060 \), and wave 4, \( b = -.426, SE = .122, p < .001 \). However, fathers’ depressive symptoms did not predict changes in mothers’ depressive symptoms at other waves. Mothers’ depressive symptoms showed no effect on fathers’ depressive symptoms across time. Similarly, youth’s time-varying internalizing symptoms did not predict changes in mothers’ and fathers’ time-varying depressive symptoms.

**Moderation Analyses.** A series of multigroup analyses tested whether the previous trait and time-varying model (Model B) varied by family ethnicity, family structure, or child gender. For each moderator, an unconstrained model was compared to a model that equated the parameter estimates across groups. Findings from the multigroup analyses revealed that the unconstrained models differed significantly from the fully equated models for family ethnicity, \( \Delta \chi^2(44) = 80.922, p < .001 \), and child gender, \( \Delta \chi^2(44) = 67.777, p = .012 \), but not family structure, \( \Delta \chi^2(44) = 48.697, p = .290 \). A series of multigroup analyses were conducted for family ethnicity and child gender to compare the unconstrained model to constrained versions of the model (e.g., trait-invariant differences, time-varying differences, mother effects on child, father effects on child, child effects on each parent). Sets of effects were tested, rather than individual paths, to avoid capitalizing on chance (i.e., type 1 error), though analyzing sets of effects may also obscure significant path coefficients (i.e., type 2 error). Significant moderation findings are reported below.

**Family ethnicity.** A chi-square difference test examining the trait-invariant differences between European and Mexican American families was significant, \( \Delta \chi^2(3) = 11.143, p = .011 \). Among European American families, youth’s trait-invariant internalizing symptoms were significantly correlated with mothers’ \( (c = .244, SE = .063, p < .001) \) and fathers’ \( (c = .238, SE = .065, p < .001) \) trait-invariant depressive symptoms, and mothers’ trait-invariant depressive
symptoms were significantly correlated with fathers’ trait-invariant depressive symptoms, \( c = .298, SE = .099, p < .001 \). In contrast, although mothers’ and fathers’ trait-invariant depressive symptoms were significantly correlated in Mexican American families, \( c = .250, SE = .119, p = .035 \), youth’s trait-invariant internalizing symptoms were not associated significantly with mothers’ \( (c = .116, SE = .070, p = .095) \) or fathers’ \( (c = -.019, SE = .050, p = .704) \) trait-invariant depressive symptoms. Other multigroup analyses did not differ significantly across European and Mexican American families.

**Child gender.** A significant chi-square difference test indicated that the time-varying effects in youth’s internalizing symptoms differed for males and females, \( \Delta \chi^2(4) = 10.580, p = .032 \). Females evidenced greater stability in their internalizing symptoms than males. The time-varying coefficients for females were significant from wave 1 to wave 2, \( b = .285, SE = .102, p = .005 \), and from wave 4 to wave 5, \( b = .825, SE = .040, p < .001 \), but not from waves 2 through 4. In contrast, the time-varying coefficients for males were significant across all waves, \( b = .359, SE = .119, p < .001 \) to \( b = .914, SE = .042, p < .001 \). The standardized coefficients for the global trait factor of internalizing symptoms were consistent with the time-varying coefficients. Specifically, the standardized coefficients for the global trait factor revealed that less of the variation in internalizing symptoms was explained by trait-invariant differences among males, with coefficients ranging from .452 to .507, than among females, with coefficients ranging from .711 to .846.

A second chi-square difference test for gender was significant, \( \Delta \chi^2(5) = 11.618, p = .040 \), revealing that the influence of mothers’ depressive symptoms on youth’s internalizing symptoms differed between males and females across time. An examination of the unstandardized path coefficients across waves indicated that mothers’ depressive symptoms at wave 4 predicted
higher levels of youth’s internalizing symptoms at wave 5 for females, $b = .063$, $SE = .024$, $p = .008$, but not for males, $b = -.002$, $SE = .016$, $p = .916$.

**Externalizing Symptoms**

Cross-lagged panel analyses evaluated the effects of mothers’ and fathers’ depressive symptoms on youth’s externalizing symptoms, as well as youth’s reciprocal influences on mothers’ and fathers’ depressive symptoms. Similar to the model for youth’s internalizing symptoms, mothers’ and fathers’ depressive symptoms at waves 1 and 4 were allowed to predict changes in youth’s externalizing symptoms at wave 5. In addition to the stability coefficients and cross-lagged influences across all three reporters, this baseline trait and time-varying cross-lagged model included within-wave covariances between all three reporters’ time-varying symptoms. Model A fit the data well, $\chi^2(44) = 64.060$, $p = .026$, RMSEA = .034 [ .012 , .051 ], CFI = .990, TLI = .975, SRMR = .035. However, a subsequent model (Model B) with only the within-wave covariances of the time-varying components for all three reporters at wave 1 also fit the data well, $\chi^2(56) = 80.945$, $p = .016$, RMSEA = .034 [ .015 , .049 ], CFI = .987, TLI = .976, SRMR = .038, and did not differ significantly from Model A, $\Delta \chi^2(12) = 16.885$, $p = .154$, which suggests that these covariances did not contribute significant information to the model.

Findings from Model B revealed both trait-invariant and time-varying differences in youth’s externalizing symptoms across time with standardized coefficients on this global trait factor ranging from .500 to .927 (see Figure 2). From waves 1 through 4, youth’s time-varying externalizing symptoms were significantly correlated with each subsequent wave, with the results showing a declining trend in externalizing symptoms across time, $b = .330$, $SE = .047$, $p < .001$ to $b = .189$, $SE = .052$, $p = .001$. Youth’s externalizing symptoms did not follow a simplex pattern (e.g., higher correlations between variables closer in time intervals), suggesting that some
effects from earlier waves did not carry through to the next wave in sequential order (see Table 2). As a result, there were significant nonadjacent paths between waves 1 and 3, \( b = .239, SE = .030, p < .001 \), and between waves 2 and wave 4, \( b = -.050, SE = .024, p = .037 \), though the latter association was negative. The time-varying association between waves 4 and 5 was not significant, \( b = .005, SE = .210, p = .980 \), which suggests that youth’s externalizing symptoms showed considerable variation from early to middle adolescence, but not from late adolescence and young adulthood. Indeed, the standardized coefficients of the global trait factor mirrored the findings of the time-varying coefficients, which showed an increase in the trait-like variation of youth’s externalizing symptoms from wave 1, \( \beta = .500, SE = .024, p < .001 \), to wave 5, \( \beta = .927, SE = .035, p < .001 \).

Findings for mothers’ and fathers’ trait-invariant and time-varying depressive symptoms were consistent with the youth internalizing model with most of the variation in mothers’ and fathers’ depressive symptoms being explained by between-person differences. An examination of the correlations between the global trait factor for all three reporters revealed that youth’s trait-invariant externalizing symptoms were marginally associated with mothers’ depressive symptoms, \( c = .171, SE = .102, p = .093 \), but were not related to fathers’ depressive symptoms, \( c = .132, SE = .089, p = .137 \). Mothers’ and fathers’ trait-invariant depressive symptoms were significantly associated, \( c = .292, SE = .077, p < .001 \).

In the total sample, mothers’ time-varying depressive symptoms at wave 1 predicted higher rates of youth’s externalizing symptoms at wave 5, \( b = .181, SE = .053, p = .001 \), but mothers’ time-varying depressive symptoms at wave 3 predicted lower rates of youth’s externalizing symptoms at wave 4, \( b = -.094, SE = .055, p = .089 \). There were no other parent effects on youth externalizing symptoms across time. Consistent with the youth’s internalizing
model for the total sample, fathers’ time-varying depressive symptoms at waves 2 and 3 were associated with lower levels of mothers’ time-varying depressive symptoms at waves 3 and 4. In addition, mothers’ time-varying depressive symptoms at wave 1 were correlated with fathers’ time-varying depressive symptoms, $c = .331, SE = .092, \, p < .001$, and youth’s time-varying externalizing symptoms, $c = .521, SE = .203, \, p = .010$, at the same time point, but youth’s time-varying externalizing symptoms at wave 1 were not correlated with father time-varying depressive symptoms, $c = .102, \, SE = .172, \, p = .556$. Youth’s time-varying externalizing symptoms at wave 1 predicted higher levels of mothers’ depressive symptoms at wave 2, $b = .081, \, SE = .034, \, p = .016$, whereas youth’s time-varying externalizing symptoms at wave 4 predicted lower levels of mothers’ depressive symptoms at wave 5, $b = -.405, \, SE = .116, \, p = .001$.

**Moderation Analyses.** A series of multigroup analyses were conducted to determine whether the previous trait and time-varying cross-lagged model (Model B) varied by family ethnicity, family structure, and child gender. For each moderation analysis, an unconstrained model was compared to a model that equated the parameter estimates across groups. Findings from the multigroup analyses revealed that the unconstrained models differed significantly from the fully equated models for family ethnicity, $\Delta \chi^2(46) = 71.498, \, p = .009$, and child gender, $\Delta \chi^2(46) = 63.835, \, p = .041$, but not family structure, $\Delta \chi^2(46) = 34.035, \, p = .904$. Several multigroup analyses were conducted for family ethnicity and child gender to compare an unconstrained model to various constrained versions of the model (e.g., trait-invariant differences, time-varying differences, mother effects on child, father effects on child, child effects on each parent). Significant moderation findings are reported below.
Family ethnicity. A chi square difference test for the time-varying covariances at wave 1 between all three reporters was significant, $\Delta \chi^2(3) = 10.126, p = .017$. Although the association between mothers’ and fathers’ time-varying depressive symptoms at wave 1 was pronounced in both European, $b = .227, SE = .126, p = .072$ and Mexican, $b = .444, SE = .142, p = .002$, American families, the relation between fathers’ depressive symptoms and youth’s externalizing symptoms differed by family ethnicity. Specifically, fathers’ depressive symptoms were significantly associated with youth’s externalizing symptoms in European American families at wave 1, $c = .541, SE = .251, p = .031$, whereas fathers’ depressive symptoms were not associated significantly with youth’s externalizing symptoms in Mexican American families, $c = -.404, SE = .255, p = .114$, at the same time point. In Mexican American families, mothers’ time-varying depressive symptoms were marginally associated with youth’s externalizing symptoms at wave 1, $c = .589, SE = .318, p = .064$, but this relation was not significant in European American families, $c = .425, SE = .265, p = .108$.

Child gender. The chi-square difference test investigating the effects of youth’s time-varying externalizing symptoms on mothers’ time-varying depressive symptoms, $\Delta \chi^2(4) = 8.226, p = .082$, and of fathers’ time-varying depressive symptoms on mothers’ time-varying depressive symptoms, $\Delta \chi^2(4) = 8.006, p = .091$, were marginally significant, therefore, these analyses were not probed further for moderation effects.

Discussion

This investigation evaluated the unique contributions of mothers’ and fathers’ depressive symptoms to youth’s internalizing and externalizing symptoms from early adolescence to young adulthood, as well as the reciprocal effects of youth’s symptomatology on each parent’s depressive symptoms. This is one of the first longitudinal studies to examine the effects of
parents’ psychopathology on youth adjustment while taking into account the influence of the co-parent’s psychopathology. Moreover, whereas prior studies have typically employed cross-sectional approaches to study young, European American children in clinical samples, this study evaluated associations between parents’ depressive symptoms and youth’s symptomatology from early adolescence to young adulthood in a community sample of European American and Mexican American families. Overall, the findings showed that both mothers’ and fathers’ between- and within-person differences in depressive symptoms across time were related to youth’s internalizing and externalizing symptoms from early adolescence to young adulthood, but reciprocal effects from youth’s symptomatology to parents’ depressive symptoms were modest. Importantly, these associations varied significantly across groups defined by family ethnicity and child gender, but not by family structure.

**Fathers’ and Mothers’ Depressive Symptoms**

Fathers’ and mothers’ depressive symptoms evidenced predominantly trait-invariant (i.e., between-person) differences, which indicates that mothers’ and fathers’ depressive symptoms remained stable over time and had comparable effects on youth’s symptomatology across early adolescence and young adulthood. Mothers’ and fathers’ between-person differences in depressive symptoms across time and their individual differences in depressive symptoms at the initial wave of assessment were positively correlated with each other, and did not significantly differ across racial/ethnic, family structure, or gender groups. Previous research has suggested similar patterns, such that, if one partner has a history of depression, there is an increased likelihood that the other partner will experience similar rates of depression (e.g., Hammen, Katz, & Le Brocque, 2002). These patterns likely reflect assortative mating patterns wherein partners both seek out individuals who are genetically similar to them and/or have similar personal
characteristics, such that, as time unfolds, fathers and mothers become increasingly psychologically similar in the context of a co-constructed environment (Watson et al., 2004).

**Youth’s Internalizing Symptoms**

Unlike mothers’ and fathers’ depressive symptoms, youth’s internalizing symptoms evidenced both between- and within-person differences from early adolescence to young adulthood. However, the between-person differences in youth’s internalizing symptoms (i.e., the differences in internalizing scores between two people) remained consistent across time, whereas the within-person differences in youth’s internalizing symptoms (i.e., the differences in an individual’s internalizing scores across time after the trait portion has been removed) became increasingly stable over time. This finding is consistent with recent work examining time-invariant between-person and time-varying within-person differences in children’s depressive symptoms, which suggests that between-person differences in children’s depressive symptoms are congruent across time, informants, and settings (Cole et al., 2017). The increasing stability of individual differences in youth’s internalizing symptoms from late adolescence to young adulthood is also consistent with previous research showing that youth’s internalizing symptoms, especially depression, increase from early to mid-adolescence, but remain stable after mid-adolescence and into adulthood (Ge, Natsuaki, & Conger, 2006). However, as in prior studies (Ge, Lorenz, Conger, Elder, & Simons, 1994; Ge et al., 2006; Kessler, 2003), girls experienced more stability than boys in their internalizing symptoms by mid adolescence and evidenced increasing rates of internalizing symptoms across early adolescence.

Between-person differences in youth’s internalizing symptoms were associated with between-person differences in mothers’ and fathers’ depressive symptoms across time. Specifically, mothers and fathers who experienced higher levels of depressive symptoms were
more likely to have children with elevated rates of anxiety and depressive symptoms from early adolescence to young adulthood. These findings are consistent with previous empirical evidence suggesting that both mothers’ and fathers’ depressive symptoms contribute to youth’s internalizing symptoms (Cummings et al., 2014; Elgar et al., 2007; Hammen et al., 2008).

Although the associations of mothers’ and fathers’ depressive symptoms with youth’s internalizing symptoms were consistent across intact and stepfather families and across families with daughters and sons, these effects varied by family ethnicity. European American youth’s experiences of anxiety and depressive symptoms were linked to both their mothers’ and fathers’ depressive symptoms, but neither set of relations attained significance among Mexican American youth. This finding contradicts prior studies suggesting that relations between parental psychopathology and child symptomatology are stronger in non-European American families because of the relatively greater interdependence among family members (Wilson & Durbin, 2010). In contrast to prior assertions, the current findings may reflect a buffering effect of close and supportive connections in Mexican American families, such that they mitigate the influence of parents’ depressive symptomatology on children.

Mirroring the obtained patterns of between-person differences, individual differences in mothers’ and fathers’ depressive symptoms predicted changes in youth’s internalizing symptoms. Specifically, mothers’ and fathers’ depressive symptoms in early adolescence (i.e., distal effects) evidenced a stronger influence on youth’s later internalizing symptoms than parents’ depressive symptoms in late adolescence (i.e., proximal effects). These findings are consistent with prior theoretical frameworks that conceptualize development as cumulative, orderly, and hierarchically integrated, such that early experiences in the family environment tend to have special significance or meaning for youth development (Cummings & Cicchetti, 1990;
Duggal, Carlson, Sroufe, & Egeland, 2001; Sroufe, 1990). In this view, distal parental depressive symptoms may lead to youth’s symptomatology in young adulthood because parents’ experiences of depressive symptoms during youth’s earlier development influence their responsiveness and psychological availability to their children’s developmental needs, which in turn, affects their children’s emergent representational, regulatory, and relational structures (Cummings & Cicchetti, 1990; Duggal et al., 2001). Alternatively, the relatively modest effect of proximal parental depressive symptoms on youth’s symptomatology may reflect the normative decline in parental influence across adolescence and into young adulthood as youth’s relationships beyond the family with platonic and romantic peers takes on increasing salience (Furman & Buhrmester, 1992; Larson, Richards, Moneta, Holmbeck, & Duckett, 1996).

The influence of mothers’ and fathers’ depressive symptoms on youth’s internalizing symptoms differed such that elevated rates of mothers’ depressive symptoms in early adolescence were associated with higher levels of youth’s internalizing symptoms in young adulthood, whereas elevated rates of fathers’ depressive symptoms in early adolescence were associated with lower levels of youth’s internalizing symptoms in young adulthood. The obtained findings are consistent with prior studies showing that mothers’ depressive symptoms in early adolescence predicted children’s anxiety and depressive symptoms in young adulthood (Reeb et al., 2015), as well as with other studies suggesting that changes in mothers’ depressive symptoms were associated with higher rates of emotional problems in children during adolescence (Cummings et al., 2014; Elgar et al., 2007). Interestingly, higher levels of fathers’ depressive symptoms in early adolescence were associated with less vulnerability to internalizing symptoms for youth in young adulthood. Given this counterintuitive finding, we encourage
efforts to replicate this pattern in advance of offering tenuous interpretations, particularly as the large number of analyses in the current study may have inflated the risk of Type 1 errors.

Importantly, although mothers’ depressive symptoms in early adolescence were positively associated with youth’s internalizing symptoms for the total sample, changes in mothers’ depressive symptoms during late adolescence emerged as a stronger influence on girls’ than on boys’ internalizing symptomatology during young adulthood. These findings are consistent with prior evidence that mothers’ depressive symptoms influence both boys’ and girls’ internalizing symptoms (Connell & Goodman, 2002), but they further suggest that girls may be especially vulnerable to mothers’ depressive symptoms during young adulthood. This pattern may reflect a shift in the relational dynamics between mothers and their children such that mothers may be more willing to share their emotional challenges with their older daughters and may depend on them for emotional support to a greater extent than their older sons. Alternately, some scholars have argued that daughters may become overinvolved with their mothers’ depression, which increases the immediate transmission of mothers’ depressive symptoms to their daughters relative to their sons (Duggal et al., 2001). Finally, changes in youth’s internalizing symptoms did not affect parents’ depressive symptoms. This pattern could suggest that youth’s internalizing symptoms create fewer disruptions in the family system, perhaps because they are more likely to go unnoticed.

Youth’s Externalizing Symptoms

Youth’s externalizing symptoms evidenced both between- and within-person differences from adolescence to young adulthood. The trait-like variation in youth’s externalizing symptoms increased from adolescence to young adulthood and the time-varying effects decreased across time. The obtained findings mirror those of prior studies, which have shown that children’s
Youth’s between-person differences in externalizing symptoms were associated with mothers’ between-person depressive symptoms, but not with fathers’ between-person depressive symptoms. Thus, mothers who had higher levels of depressive symptoms were more likely to have children who engaged in delinquent, intrusive, and rule-breaking behaviors from early adolescence through young adulthood. Although these findings support prior studies showing that depressed mothers tend to have children with more problem behaviors (Elgar et al., 2007; Natsuaki et al., 2014), the current results are inconsistent with previous studies showing that fathers’ depressive symptoms are more important than mothers’ depressive symptoms for understanding children’s externalizing symptoms (Connell & Goodman, 2002; Kane & Garber, 2004; Low & Stocker, 2005). Given that prior studies have typically examined mothers’ and fathers’ depressive symptoms in separate models, and also have not disentangled between-person from the within-person effects, the association between fathers’ depressive symptoms and youth’s externalizing symptoms in the current study may have been attenuated by the concomitant consideration of mothers’ symptomatology, as well as the separation of these two variance types in this study.

Similar to the influence of parent’s between-person differences in depressive symptoms on youth’s externalizing symptoms, within-person differences in mothers’, but not fathers’, depressive symptoms predicted changes in youth’s externalizing symptoms. Interestingly, whereas mothers’ elevated depressive symptoms during early adolescence (i.e., distal effects) predicted higher levels of youth’s externalizing symptoms during young adulthood, mothers’
elevated depressive symptoms in middle adolescence (i.e., proximal effects) predicted lower levels of youth’s externalizing symptoms during late adolescence. Although the distal effect found for mothers’ depressive symptoms is consistent with prior research (Cummings et al., 2014; Elgar et al., 2007), the proximal negative effect of mothers’ depressive symptoms on youth’s externalizing problems is inconsistent with the research literature and warrants further replication in the field before it should be interpreted.

The current findings also highlight the importance of investigating child effects on parents’ psychological functioning and suggest that children’s behavior in adolescence and young adulthood may feed back to influence parents’ mental health in divergent ways. Youth’s externalizing symptoms predicted higher levels of mothers’ depressive symptoms during early adolescence, whereas youth’s externalizing symptoms in late adolescence predicted lower levels of mothers’ depressive symptoms during young adulthood. These data suggest that youth’s externalizing problems during early adolescence were associated with elevated depressive symptoms in mothers, but these same problems were associated with lower maternal depressive symptoms in late adolescence. It may be that mothers are more likely to attribute their children’s problem behaviors during early adolescence to their own shortcomings as parents, whereas higher levels of youth’s externalizing behaviors during late adolescence might be perceived as more age-appropriate or normative.

**Strengths, Limitations, and Future Directions**

This investigation drew on a multi-wave, multi-informant longitudinal study of a large and diverse sample of two-parent families. Notable strengths of this study included the examination of between- and within-person effects, transactional effects among multiple family members, proximal and distal influences, and the ability to evaluate directional hypotheses and
UNIQUE EFFECTS OF MOTHERS’ AND FATHERS’ DEPRESSIVE SYMPTOMS

developmental timing. However, several limitations both qualify the interpretation of the findings and reveal promising directions for future research.

First, given the complexity of the models and moderate sample size, it was not possible to test synergistic effects between the independent and contextual variables in this study. Although some researchers have argued that the effects of parental depression on youth adjustment may be additive (Reeb et al., 2010; Reeb et al., 2015), rather than interactive, previous studies have found interactive effects between mothers’ and fathers’ depressive symptoms on youth adjustment (Brennan et al., 2002). Furthermore, it is important to note that higher level interactions among family ethnicity, family structure, and child gender could lead to differential developmental outcomes. Likewise, we encourage future researchers to consider alternative analytic models, such as dynamic growth curve models (McArdle & Nesselroade, 2003).

Second, moderation analyses were tested in sets (e.g., all mother effects on youth’s symptomatology across time) to mitigate type 1 error. However, this analytic approach could have masked significant effects for individual paths. In addition, the complexity of the models as well as the sample size may not have been sufficient to detect small effects. Therefore, future research should utilize larger, nationally representative datasets to determine if the effects found in this study are consistent across different samples. Future research should also determine whether the distal effect of fathers’ depressive symptoms on youth’s internalizing symptoms differs between sons and daughters.

Third, we were not able to disentangle the confound of family structure and genetic influences on youth’s symptomatology. Indeed, a recent study examining the interplay between genetics and early environmental influences on youth’s internalizing symptoms from late childhood to adolescence found that genetic predisposition was the sole predictor of the stable
latent trait (i.e., time-invariant) portion of youth’s internalizing symptoms across adolescence (Musci et al., 2016). However, although a fully unconstrained model by family structure and models with only fathers showed a direct association between fathers’ depressive symptoms and youth’s internalizing symptoms, this study did not provide strong support for paternal genetic transmission as family structure was not found to moderate the obtained pathways.

Finally, the factor score estimates for youth’s internalizing and externalizing symptoms were computed using a novel approach to data harmonization and this process may not have been fully corrective. For example, the associations between waves 3 and 4 were lower (and in some instances not significant), and this was coincident with the inter-wave transition in measurement tools. That said, it is important to note that the time-varying associations between these two waves were quite large considering this was the longest time interval in the study. Although the findings were consistent with prior research on youth’s internalizing (Cole et al., 2017; Ge et al., 2006) and externalizing (Galambos et al., 2003; Kim et al., 2003) symptoms, future research should verify these trait and time-varying differences in youth’s symptomatology in studies with closer assessment intervals. It is also important to note that although we were able to establish weak invariance with youth’s measures of internalizing and externalizing symptoms across time, we were unable to acquire strong invariance which is fundamental in assessing change across time. Therefore, the time-varying effects could be attributed to both real change as well as measurement error and should be interpreted with caution.

Implications and Conclusions

The current study addressed several gaps in the existing literature on the effects of parent’s depressive symptoms on youth’s adjustment. Overall, the findings revealed dynamic transactions across family members and revealed opportune times and targets for intervention
and prevention efforts aimed at mitigating the deleterious effects of parental psychopathology on adolescent and young adult adjustment. In particular, these data suggest that intervention and policy efforts to reduce the negative impact of parental psychopathology on children should include the whole family system. Furthermore, the findings from this study highlight the need to include children in intervention efforts as youth’s externalizing symptoms influenced parents’ depressive symptoms.

One of the most interesting findings in this study was the moderating effect of family ethnicity on the association between parental depressive symptoms and youth’s symptomatology, with European American youth appearing more susceptible to their parents’ depressive symptoms than their Mexican American peers. In order to maximize the effectiveness of prevention and intervention programs, it might be beneficial for researchers and clinicians to understand what features of the Mexican American family environment seem to prevent the transmission of negative parental depression effects on youth adjustment. Intervention efforts also need to consider developmental timing during the implementation of these programs.

Finally, given the unique examination of trait and time-varying effects in this study, some of the findings were inconsistent with previous empirical evidence. In particular, most of the findings that were consistent with previous literature were at the trait level (i.e., between-person differences), suggesting that findings in the existing literature on longitudinal and transactional effects are driven in large part by differences at the trait level and less so by time-varying effects. This investigation suggests that researchers need to be careful about the conclusions they draw from traditional cross-lagged models and should adhere to recent calls in the field (Berry & Willoughby, 2016; Hamaker et al., 2015) that advocate for the separation of between- and within-person differences in psychological constructs across time.
References


### Table 1. Correlations, Means, and Standard Deviations for Mothers’ and Fathers’ Depressive Symptoms

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| M         | 4.49| 4.55| 4.56| 4.76| 4.49| 4.22| 4.17| 4.20| 4.33| 4.16 |
| SD        | 1.62| 1.77| 1.60| 1.75| 1.78| 1.45| 1.32| 1.40| 1.53| 1.37 |

Note. MDEP = Mothers’ Depressive Symptoms; FDEP = Fathers’ Depressive Symptoms. Correlations equal to or greater than .099 are significant at the probability level of .05. Correlations equal to or greater than .130 are significant at the probability level of .01.
Table 2. Correlations for the Latent Constructs of Youth’s Externalizing Symptoms Across Waves

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Figure 1. Unstandardized coefficients for the trait and time-varying cross-lagged panel analysis for youth’s internalizing symptoms. *p ≤ .05. **p ≤ .01.
Figure 2. Unstandardized coefficients for the trait and time-varying cross-lagged panel analysis for youth’s externalizing symptoms. *$p \leq .05$. **$p \leq .01$. 