Ablation of Leptin Signaling to Somatotropes: Changes in Metabolic Factors that Cause Obesity

Noor Akhter, Angela K. Odle, Melody L. Allensworth-James, Anessa C. Haney, Mohsin M. Syed, Michael A. Cozart, Streamson Chua, Rhonda Kineman, and Gwen V. Childs

Department of Neurobiology and Developmental Sciences (N.A., A.K.O., M.L.A.-J., A.C.H., M.S.S., M.A.C., G.V.C.), College of Medicine, University of Arkansas for Medical Sciences, Little Rock, Arkansas 72205; Department of Medicine (S.C.), Albert Einstein College Medicine, Bronx, New York; and Jesse Brown Veteran Affairs Medical Center (R.K.), Research and Development Division and Department of Medicine, University of Illinois at Chicago, Chicago, Illinois 60612

Mice with somatotrope-specific deletion of the Janus kinase binding site in leptin receptors are GH deficient as young adults and become obese by 6 months of age. This study focused on the metabolic status of young (3–4.5 month old) preobese mutant mice. These mutants had normal body weights, lean body mass, serum leptin, glucose, and triglycerides. Mutant males and females showed significantly higher respiratory quotients (RQ) and lower energy output, resulting from a higher volume of CO₂ output and lower volume of O₂ consumption. Deletion mutant females were significantly less active than controls; they had higher levels of total serum ghrelin and ate more food. Mutant females also had lower serum insulin and higher glucagon. In contrast, deletion mutant males were not hyperphagic, but they were more active and spent less time sleeping. Adiponectin and resistin, both products of adipocytes, were increased in male and female mutant mice. In addition, mutant males showed an increase in circulating levels of the potent lipogenic hormone, glucose-dependent insulinotropic peptide. Taken together, these results indicate that mutant mice may become obese due to a reduction in lipid oxidation and energy expenditure. This may stem from GH deficiency. Reduced fat oxidation and enhanced insulin sensitivity (in females) are directly related to GH deficiency in mutant mice because GH has been shown by others to increase insulin sensitivity and fat oxidation and reduce carbohydrate oxidation. Gender-dependent alterations in metabolic signals may further exacerbate the future obese phenotype and affect the timing of its onset. Females show a delay in onset of obesity, perhaps because of their low serum insulin, which is lipogenic, whereas young males already have higher levels of the lipogenic hormone, glucose-dependent insulinotropic peptide. These findings signify that leptin signals to somatotropes are vital for the normal metabolic activity needed to optimize body composition. (Endocrinology 153: 4705–4715, 2012)

Leptin, a product of white fat cells (1), is an important cytokine that is best known for its regulation of appetite and food intake by way of specific receptors (LEPR) in the hypothalamic neurons. Evidence, based on both in vivo and in vitro studies, has shown that leptin also regulates other cell types, including pituitary somatotropes. Animals, which cannot make leptin (2, 3), show reduced numbers of somatotropes (2–5). Humans with impaired leptin signaling, due to a mutation in LEPRb (exon 16), have impaired growth (6).

Leptin receptors are expressed in the majority of somatotropes in rats (7, 8) and sheep (9) or pigs (10), and Cai and Hyde (11) reported increased expression of LEPRb in pituitaries of mice with somatotrope hyperplasia [human GHRH transgenic mice (12)]. Our recent studies of both rats and mice agree that most somatotropes bear leptin receptors (13–15).
However, the importance of leptin to somatotrope function is still not well understood. The evidence showing that leptin deficiency reduced the numbers of somatotropes (16, 17) was questioned because the reduction could have been caused by the metabolic disease itself or the hypogonadal condition (18–20). Luque et al. (21) reported that leptin restored GH secretion and increased pituitary GHRH receptors in ob/ob mice that were infused with exogenous leptin for 7 d, which could not be attributed to changes in hypothalamic GHRH expression. Our recent in vitro studies reported that exposure to 10–100 pg/ml leptin increased the percentages of somatotropes reduced by 24 h of fasting, and 1–10 pg/ml of leptin stimulated an increase in GH mRNA in normal pituitary somatotropes (14). Collectively these observations pointed to the importance of direct interactions of leptin on somatotropes.

To better understand the impact of leptin signaling on somatotropes, we recently used a selective knockout strategy to ablate the signaling portion of leptin receptors in pituitary somatotropes (22). Mice bearing Cre-recombinase driven by the rat growth hormone promoter (rGHp-Cre) (23) were crossed with mice bearing leptin receptor gene alleles in which exon 17 was flanked by loxP (Lepr exon 17 flox/flox) (24, 25). Somatotrope-specific loss in leptin signaling reduced the population of immunolabeled somatotropes and serum GH by 60%. The GH deficiency (GHD) was detected as early as 3 months of age, but body weights did not significantly differ between mutant and control mice until 5 months in males and 6.5 months in females. dual-energy x-ray absorptiometry analysis of the older animals showed a clear increase in fat mass, with no change in lean mass. Serum leptin levels were normal, although they increased appropriately with increasing fat mass.

The delay in fat mass accumulation in the mutant mice, presents the opportunity to identify behaviors and factors that might eventually drive the changes in body composition, observed in later life. To this end, a comprehensive laboratory animal monitoring system (CLAMS) was used to assess activity level, sleep episodes, and food/water intake as well as O2 consumption and CO2 output by indirect calorimetry. These data were correlated with changes in circulating adipokines and proinflammatory cytokines known to be associated with obesity.

Materials and Methods

Production of deletion mutant mice lacking LEPR exon 17 in somatotropes

Breeding cages were set up to produce N6 and N7 generation deletion mutants and littermate controls (FVB/NJ) as previously described (22). Mice were housed five per cage and maintained at 10-h light,14-h dark cycle until they were old enough for testing (3–4.5 months of age). The animals were fed Teklad 8640 or Teklad 22/5 rodent diet (Tekland, Madison, WI), which is 22% protein, 5% fat, and 4.5% fiber. While breeding, the mice were fed Teklad Global 19% protein diet 2019, which is 19% protein, 9% fat, and 5% fiber. The Animal Care and Use Committee approved all protocols, annually (protocol 3014).

Comprehensive laboratory animal monitoring system

The indirect calorimetry, activity levels, and food intake were determined using the Oxymax CLAMS (Columbus Instruments, Columbus, OH) over a period of 72 h. This unit is in the same room that houses the breeding colonies, which facilitates acclimation. Animals were acclimated in the CLAMS unit for 48 h before the 72-h test period and received water and food (powdered Teklad 8604 diet). A total of four different groups (four animals per group) of male or female mutant and littermate control mice were evaluated. Within and between groups, they were age and weight matched. For example, in a given experiment, if we had a 3-month-old and a 4-month-old deletion mutant, we matched this with a 3-month-old and a 4-month-old control. For each animal, we calculated lean body mass, which was body weight (M) to the 3⁄4 power, according to Kleiber’s law (26, 27). This formula is a well-established surrogate for metabolic mass. Their weight was loaded into the CLAMS program and the conversion to lean body mass was built into the software used by the CLAMS, which normalized the volume of oxygen, carbon dioxide, and the resulting respiratory quotient (RQ) to lean body mass.

The activity monitors in CLAMS also are an indirect means of detecting sleep. A sleep epoch was defined as a 60-sec bout of no activity. The computer-generated sleep analysis recorded the number of consecutive sleep epochs and the total percent of time sleeping for each mouse. This system has been shown to give excellent agreement with other sleep detection approaches, including electromyographic/electroencephalographic (EEG) and video monitoring (28).

The output from the software analysis program (CLAX, Columbus Instruments) was analyzed using Instat3 or PRISM 5.0 software (both from GraphPad Inc., San Diego, CA). One-way ANOVA was run to detect significant differences between mutant and control mice within gender. Bonferroni and Neuman Keuls multiple comparisons tests were used as post hoc tests. The percent relative cumulative frequency approach (29) was used to determine the frequency of RQ values. The differences between mutant and control values were assessed by nonlinear regression curve fitting and Mann Whitney and D’Agostino and Pearson omnibus normality tests.

Multiplex enzyme immunoassays for cytokines and adipokines

Trunk blood samples were collected from the mice at the time the animals were killed between 0900 and 1100 h to avoid diurnal variations. The animals were 3.5–4 months of age. In addition, serum from 20–30 additional age and weight-matched mice were taken at the same time of day. These mice were littermates or from litters born during the same time period. This group provided serum from animals that had not been run in the CLAMS to determine differences that might have come from having been in the CLAMS environment. No differences were detected in their weights or in the values themselves. Therefore,
the serum values are averaged with those from mice exposed to the CLAMS. All serum samples were assayed in the Luminex LX200 xPONENT 3.1 with Millipore Multiplex adipocyte, adipokine, or metabolic disease panels designed for mouse analytes (Millipore, Bedford, MA). The analytes included leptin, ghrelin, adiponectin, resistin, glucose insulino-tropic peptide (GIP), IL-6, insulin, glucagon, plasminogen activator inhibitor-1 (PAI-1), monocyte chemotactic protein-1 (MCP-1), and TNF-α. Sera were also tested for glucose with a glucometer and triglycerides with the Cayman Chemical colorimetric assay kit (Ann Arbor, MI).

Values for male or female mice were averaged and subjected to one-way ANOVA, followed by Tukey’s, Neuman-Keul’s, and Bonferroni’s multiple comparison post hoc tests. In some cases, a Student’s t test with Welch’s correction was used to compare values from control and deletion mutant animals (P < 0.05 was considered significant).

Results

Young deletion mutant mice are not obese and have normal leptin levels

As shown in Fig. 1, total mean body weights, lean body mass (Fig. 1A), and circulating leptin (Fig. 1B) did not differ between 3- and 4.5-month-old mutant mice within a given sex. The average leptin levels in females were significantly lower than that in males. With respect to food intake, the system reported the cumulative amount of food eaten over the 72-h period, with the option of reporting cumulative food eaten in only the two to three light or dark periods during that time. The system also reported the amount of food eaten at each feeding bout.

Consistent with previous reports, male and female mice consumed more food during the dark phase, independent of genotype (Fig. 1, C and D). In addition, deletion mutant males showed no differences in the cumulated grams of food eaten during dark or light phases (Fig. 1F). In contrast, the cumulated amount of food eaten by the deletion mutant females was higher than that eaten by the controls during both the dark and light phases (Fig. 1D), and mutant females ate more food/feeding bout during the dark phase (Fig. 1E). Half of the females were 3–3.5 months old and the others were 4.5 months old. When the data were separated by age, the higher food consumption was sig-

![Image of results](image-url)
significantly different in the population of older females (4.5 months, data not shown) but not in the 3-month-old group. Mutant females also showed a significant increase in circulating total ghrelin levels, which correlated with their increased food consumption (Fig. 1F).

Preobese deletion mutants have a higher respiratory quotient and lower energy expenditure

Indirect calorimetry for the four groups of males and females (Fig. 2, A–F) showed that deletion mutant females had higher RQ, especially during the nighttime feeding cycle with females. C and E, Higher EC_{50} of individual RQ values is shown on the percentile curve after the use of the percent relative cumulative frequency approach (D’Agostino and Pearson omnibus normality test \( P < 0.0001 \); Mann-Whitney test, \( P = 0.01 \), males, and \( P = 0.024 \), females). Statistics in the remaining figures were done with the \( t \) test with Welch’s correction and the InSTAT3 software (GraphPad). D and F, There is higher RQ in both night and day cycles (\( P < 0.0001 \)). G and I, The higher RQ is due to higher VCO\(_2\)/lean body mass, averaged for light and dark periods (\( P < 0.0001 \)). Figures 2 h and 2j, H and J, The higher RQ is also associated with a lower VO\(_2\), averaged for light and dark periods (\( P < 0.0001 \)). K and L, Illustration of the lower energy output averaged during light and dark phases (\( P < 0.0001 \)). Closed star, Significantly greater than control values; open star, significantly different from values in light phase.

Preobese mutant females show reduced activity and males have reduced sleep

When activity was measured by infrared beam breaks, both control and mutant female mice showed enhanced activity during the dark phase (Fig 3, A–C). However, deletion mutant females showed reduced total activity, compared with female controls, which included both ambulation and grooming (total activity, Fig. 3A), walking (Fig. 3B), and rearing or jumping (Fig. 3C). In contrast, deletion mutant males showed no significant difference in number of beam counts when the total activity in the x-axis was detected (data not shown). However, there was a significant increase in ambulatory (Fig. 3D) or rearing or
jumping activity (Fig. 3E). The sleep analysis, based on absence of activity, showed that deletion mutant males spent less time sleeping than littermate controls (Fig. 3F).

**Glucose regulation and triglycerides**

Glucose-regulatory hormones (insulin and glucagon) and C-peptide, a proinsulin, were assayed in sera from animals collected after the CLAMS experiments as well as from additional animals collected during this period. All animals had glucose levels that were within the normal ranges for young adult mice (females-controls 137 ± 110 mg/dl; mutants, 153.5 ± 9 mg/dl; males-controls, 165 ± 6 mg/dl; mutants, 173 ± 29 mg/dl). There was no difference between controls and mutants. Similarly, there were no differences in triglyceride levels between controls (males, 249 ± 54 mg/dl; females 296 ± 35 mg/dl) and mutants (males, 257 ± 114 mg/dl; females, 225 ± 25 mg/dl). The triglyceride values for both groups are slightly higher, but within the range reported for FVB strains at the Jackson Laboratory Mouse Phenome Database (http://jaxmice.jax.org/support/phenotyping/FVBNJdata001800.pdf).

Insulin was assayed with adipokine and metabolic disease panels. Figure 4A shows that there was no significant difference in serum insulin levels in the males. In contrast, deletion mutant females had significantly lower insulin levels when compared with the male populations. Furthermore, deletion mutant females showed a significantly lower level of insulin by Student’s t test when compared with control females (P < 0.02). Levels of the proinsulin, C-peptide, followed insulin levels. C-peptide levels in females were 53% of those in the control and significantly lower than in all other populations (Fig. 4B). There were also no significant differences in glucagon levels in the two groups of males. In contrast, deletion mutant females showed 1.6-fold higher glucagon levels (Fig. 4C).

**Deletion mutant mice have higher levels of adipokines, whereas alteration in inflammatory cytokines are sexually dimorphic**

Circulating adiponectin and resistin levels were significantly elevated in both male and female mutant mice compared with controls (Fig. 5, A and B). Sexually dimorphic changes were observed for circulating GIP and IL-6 in mutant mice. Specifically, GIP was elevated in males, whereas IL-6 was reduced in females (Fig. 5, C and D). No differences were observed between mutants and controls for circulating PAI-1, TNF-α, and MCP-1 (data not shown).

**Discussion**

Our laboratory has previously reported that somatotrope-specific, Cre-recombinase-mediated excision of LEPRb
exon 17, which codes for the portion of the molecule in the cytoplasmic tail that contains the Janus kinase binding site (22), blocks phosphorylation of signal transducer and activator of transcription 3 in response to leptin stimulation (22). Lack of leptin-mediated Janus kinase/signal transducer and activator of transcription signaling in somatotropes resulted in mice that grow normally but are GHD as adults.

As in cases of human adult-onset GHD (30–32), the deletion mutant mice became obese later in adult life. In the current study, we examined the metabolic profile of 3- to 4.5-month-old mutant mice before they became obese as indicated by their normal body weights and leptin levels. The objective was to determine whether the young, preobese mutants would exhibit metabolic changes and behaviors that could eventually lead to obesity. These preobese mutants showed no changes in serum glucose or triglycerides. We also observed that male mutant mice were not hyperphagic, which is consistent with their normal leptin levels and our previous report showing that leptin receptors in the hypothalamus remained intact (22). In contrast, mutant female mice ate more than controls. Whereas their leptin levels were also not different from controls, they did have significantly higher ghrelin levels.

The following discussion will point to differences in metabolic activity, behaviors, and hormones, which may predispose the onset of obesity in these animals, highlighting sex differences where appropriate.

**Mutant males and females burn less fat and more carbohydrates**

The strongest predisposition to obesity was seen in the calculation of respiratory quotients \((RQ = VCO_2 / VO_2)\) for these mutants. Indirect calorimetry demonstrated that deletion mutant males and females had higher RQ than the littermate controls. In relation to the amounts of CO\(_2\) produced, the oxidation of fat requires more oxygen than the oxidation of carbohydrates; therefore, a higher RQ indicates that animals are oxidizing carbohydrates rather than fat (33). This, in the context of their reduced energy expenditure, would lead to excess fat accumulation. These changes may be directly caused by their GHD (22). In previous reports, male mice with adult-onset isolated GHD had higher RQ values, particularly during the dark cycle (34), and were fatter than littermate controls. In humans, an increased RQ is both predictive of obesity (35–37) and associated with the obesity-inducing effects of ghrelin (38). The association between GH deficiency and fat vs. carbohydrate oxidation was shown in studies of normal and GHD humans. In the first study, GH treatment of normal humans and tissues resulted in a significant reduction in RQ and a 29% increase in fat oxidation by muscle mitochondria coupled with a 69% decrease in carbohydrate oxidation (39). Similarly, GH treatment of GHD subjects resulted in a significant decrease in hepatic carbohydrate oxidation (40). The higher RQ indicating preferential burning of carbohydrates is thus consistent with the GHD of the mutant animals in the present study. Future studies would be needed to evaluate the fat and carbohydrate oxidation in muscle and liver of these animals.

**GHD and changes in pancreatic hormones**

Whereas obesity promotes insulin resistance and dysregulation of glucose transport, GHD is often associated with increased insulin sensitivity (34). As reported in our previous studies, the preobese male and female
deletion mutant mice had normal levels of serum glucose. To determine any changes in pancreatic hormones in these animals, we assayed glucagon, insulin, and the insulin prohormone C-peptide. The preobese deletion mutant males showed no changes in any of these hormones. However, all three hormones were significantly changed in deletion mutant females. Both insulin and C-peptide were significantly reduced, whereas glucagon was significantly increased. These changes may reflect increased insulin sensitivity as a result of the GH deficiency (34). In addition, their lower insulin may explain why the female deletion mutant mice have a delay in the onset of obesity compared with the male deletion mutant mice because insulin is a potent lipogenic hormone (41, 42).

GHD and reduced activity or sleep as risk factors for obesity

GHD in humans is also associated with loss of lean body mass and muscle strength, which can be restored by GH therapy (43–45). In the present model, the loss of muscle strength or mass may be the basis behind the reduced activity seen in the deletion mutant females. The reduced activity levels also correlate with the higher ghrelin seen in these females (Fig. 1C) because recent studies have shown that ghrelin is involved in reducing spontaneous physical activity (46). Another contributing factor in females may be the reduction in IL-6, which is secreted by adipose tissue, immune cells, fibroblasts, and endothelial cells (47). Recent studies by Faldt et al. (35) have reported that IL-6 deficiency was associated with a higher RQ and a reduced endurance in young, preobese animals. The investigators suggested that endogenous IL-6 is important for high oxygen consumption that allows the animal to maintain skeletal muscle work during prolonged exercise. The lower IL-6 in the deletion mutant females thus also correlates with their reduced activity and high RQ.

In contrast, preobese male deletion mutants remain active and in fact exhibit a slight increase in activity levels over control males. This is correlated with a significant reduction in the percent of time sleeping, which may be another response to GHD. Sleep and GH secretion are related as a burst of GH secretion is seen during deep non-rapid eye movement (REM) sleep in humans and other animals (48–51). However, GH itself may be more directly involved in REM sleep. Obal et al. (52–54) reported sleep alterations in the dwarf rat (dw/dw) or mouse (lit/lit), which has defective GHRH signaling. They were able to separate regulatory pathways and reported that non-REM sleep was mediated centrally via the GHRH receptor, and REM sleep was regulated via GHRH-GH pathways. There is also significant association between sleep loss and onset of obesity and type 2 diabetes (55–58). Thus, reduced sleep could be an important factor that leads to the obesity seen in the older (5–6 months old) males (22), although it is clear that the quality of sleep may need to be monitored by EEG recordings to...
fully evaluate its impact. It is worthwhile to note that Pack et al. (28) reported an 88–94% agreement between EEG recordings, videography, and the activity readings in the Oxymax CLAMS unit in mice. Their study showed that the mean difference per 2-h interval between inactivity-defined sleep (used in our study) and EEG/electromyogram defined sleep was only 1 min.

**Adipokine secretion as risk factors in preobese mutants**

Adiponectin is secreted by adipocytes and increases with adipogenesis (59). However, it is also secreted in inverse proportion to the body mass index (60–62). It has antidiabetic, antiatherogenic, and antiinflammatory properties and is thus protective against the damaging inflammatory effects of obesity (63, 64). Both male and female deletion mutant mice had higher adiponectin levels than controls (Fig. 4A). And in both sets of females, adiponectin levels were higher than those in males. This may be associated with adipogenesis, signifying that the animals are beginning to build excess fat stores. In addition, plasma adiponectin is inversely correlated with cytokines IL-6 and TNF-α (47), and deletion mutant females do show reduced IL-6 (Fig. 4D).

Resistin is a cytokine that is also expressed primarily by adipocytes and is positively correlated with fat mass. Its function is to antagonize insulin action by inhibiting glucose uptake in skeletal muscle and impairing insulin actions on hepatic glucose production (65–67). It may be an important risk factor for the development of insulin resistance and, therefore, we reasoned that animals might show higher resistin levels. In the present studies, the preobese deletion mutant males did show higher resistin levels than controls by Student’s *t* test. Mice with developmental GHD or GH resistance as well as adult-onset isolated GHD mice have been shown to be more insulin sensitive relative to controls (34), consistent with the fact that GH, at least at high levels, reduces insulin sensitivity in humans (68–70) and mice (71). Future studies will be needed to correlate their higher resistin with changes in insulin resistance in these animals, especially after they become obese.

GIP, also known as the glucose-dependent insulino-tropic peptide, is a 42-amino acid peptide synthesized by K cells in the mucosa of the duodenum and the jejunum (72, 73). GIP is clearly elevated in obese humans and animals (74). There are functional GIP receptors on adipocytes and GIP has been shown to increase fatty acid synthesis (75), stimulate lipoprotein lipase activity (76), and stimulate glucose transport into adipocytes and intestinal epithelial cells (77, 78). All of these actions may predispose an animal to obesity and GIP is thus a target for therapeutic modulation of activity via vaccination or receptor modulation (73, 74). In the present study, GIP levels were significantly higher in deletion mutant males than in controls. This may be seen as a risk factor for males and the fact that it appears elevated before they become obese may explain the earlier onset of obesity in the deletion mutant males.

**Inflammatory responses are not evident in preobese mutants**

Finally, because obesity is associated with low-grade inflammatory changes (79), we also tested proinflammatory cytokines to determine whether a rise in their levels might predict the eventual onset of increased fat mass. These included PAI-1, a serine protease inhibitor that is involved in angiogenesis and atherogenesis; TNF-α, a proinflammatory cytokine secreted by macrophages (80); and adipocytokines (81) and MCP-1, a chemokine that recruits monocytes to sites of inflammation (79). There were no significant increases in serum levels of PAI-1, MCP-1, and TNF-α.

**Mechanisms behind the GHD**

Our previous study showed that the GHD was associated with a lower number of cells immunolabeled for GH (22), which pointed to the importance of leptin signals needed to maintain an optimal number of somatotropes. However, more recent studies from our laboratory have discovered that the full somatotrope population in these mutants can be detected by their expression of GH mRNA (82), suggesting that the leptin-mediated optimization functions at the level of protein translation or storage. Furthermore, we discovered that the GH stores and secretion in this population can be fully restored if one exposes them, *in vitro* to ghrelin and GHRH (82). These findings suggest roles for leptin in pathways that optimize GH translation or storage, and future studies will be needed to explore these roles.

**Summary and conclusions**

We originally hypothesized that, without leptin signals, somatotropes are unable to respond appropriately to increasing adiposity by secreting sufficient GH, which is needed to maintain optimal levels of fat stores (22) by both lipolytic activity and preferential oxidation of fat. Our more recent studies (82) suggested that this may be due to defects in pathways leading to GH protein translation or storage. Whatever the mechanism causing GHD, we postulated that, in these GHD mice, conditions that prevent adiposity are left unchecked. Indirect calorimetry was run to identify metabolic and behavioral factors in preobese animals that might eventually cause fat accumulation. The
preobese animals showed increased RQ, which is a significant risk factor for obesity. In addition, there were differences in patterns of behavior and hormone levels when males and females were compared. These differences (high GIP in males and low insulin in females) may actually explain the differences in the timing of onset of obesity in this group of mutants. Future studies are needed to determine the impact of these changes on insulin resistance and glucose tolerance. It is clear from these studies that the normal signaling circuitry involving leptin from adipocytes to somatotropes is vital to help balance the metabolic activity needed to optimize body composition.

Acknowledgments

Address all correspondence and requests for reprints to: Gwen V. Childs, Ph.D., Department of Neurobiology and Developmental Sciences, University of Arkansas for Medical Sciences, College of Medicine, 4301 West Markham, Slot 510, Little Rock, Arkansas 72205. E-mail: childsgwen@uams.edu.

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