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The myometrium undergoes substantial remodeling at the time of labor including rearrangement of the cellular contractile machinery. The regulation of this process in human myometrium at the time of labor is poorly defined, but evidence in other muscle types suggests modulation by small heat shock proteins (sHSP). The aim of this study was to investigate whether similar changes in sHSP occur in the myometrium at labor. Using a quantitative proteomic approach (two-dimensional difference gel electrophoresis), we found a 69% decrease in the sHSP αB-crystallin in the myometrium at labor plus multiple isoforms of HSP27. Immunoblotting using phosphospecific HSP27 antibodies (HSP27-serine15, -78, and -82) detected marked changes in HSP27 phosphorylation at labor. Although total HSP27 levels were unchanged, HSP27-Ser15 was 3-fold higher at labor. Coimmunoprecipitation studies showed that HSP27 coprecipitates with αB-crystallin and also smooth muscle α-actin. Immunofluorescence studies demonstrated a relocation of HSP27 from the perinuclear region to the actin cytoskeleton at labor. The functional significance of these changes was demonstrated in vitro where myometrial strips stimulated to contract with oxytocin exhibited increased HSP27-Ser15 phosphorylation. Our findings provide data consistent with a novel pathway regulating human myometrial contraction at labor and identify HSP27 and αB-crystallin as potential targets for future tocolytic design. (Endocrinology 149: 245–252, 2008)

A CRITICAL ASPECT of normal human birth is the ability of uterine smooth muscle (myometrium) to respond to contractile stimulants (uterotonic) and to produce coordinated, forceful contractions. This involves a significant remodeling process that includes increased connectivity (e.g. the formation of gap junctions) between myocytes and changes in myocyte excitability as well as major changes to the cytoskeleton (1, 2).

Myometrial contractility is considered to be regulated primarily by intracellular free calcium ion concentration ([Ca\(^{2+}\)]). Increased [Ca\(^{2+}\)] results in enhanced interaction between calmodulin and myosin light chain kinase (MLCK). Phosphorylation of MLC triggers cross-bridge cycling along fibrillar actin filaments and the development of force necessary for contraction (1, 3). Relaxation of the muscle is thought to be a reversal of this process; that is, the removal of [Ca\(^{2+}\)] induces dissociation of calmodulin from MLCK. When myosin is dephosphorylated by MLCK phosphatases, it dissociates from the actin filament, and relaxation occurs. Despite widespread acceptance of this model, discrepancies remain. Smooth muscle contraction can be sustained in the continued presence of a contractile agonist, although increases in [Ca\(^{2+}\)], and MLC phosphorylation are transient (4, 5). This suggests that there are additional mechanisms involved in the regulation of smooth muscle contractility.

Changes in the cytoskeletal machinery of smooth muscle cells endow the cells with the structural and functional capacity to generate force and contract (6). The regulation of this process in the human myometrium is unknown, but data from other muscle cells suggest that small heat shock proteins (sHSP), such as αB-crystallin and HSP27, play an important role. αB-crystallin and HSP27 are highly expressed in muscle and can directly interact with each other to form homo-oligomers as well as hetero-oligomeric complexes (7–10). Furthermore, these proteins can interact with components of the actin cytoskeleton to modulate actin filament formation and thus contraction (11, 12). For example, HSP27 is a known actin-binding protein that can modulate actin filament dynamics through actin polymerization inhibition (13, 14). Additionally, overexpression of HSP27 in fibroblast cells promotes the accumulation of fibrillar actin (F-actin) (15).

The functional characteristics of sHSP are predominantly determined by phosphorylation, which induces a change in the tertiary structure of these proteins, thereby altering their affinity to form homo- or hetero-oligomers or to interact with the actin cytoskeleton. For example, phosphorylation induces the dissociation of αB-crystallin (16) and HSP27 multimers (17, 18) into small monomer and dimer subunits. HSP27 has three predominant sites of phosphorylation, serine residues 15, 78, and 82 (19, 20). The substitution of these serine residues with glycine prevents the dissociation of HSP27 oligomers and prohibits HSP27-mediated accumulation of F-actin (21), thereby supporting a role for HSP27.
phosphorylation in actin filament stabilization. Moreover, it has been proposed that the phosphorylation of HSP27 on serine residue 15 increases its propensity to bind to and stabilize actin filaments, in turn promoting contraction (11, 18).

In this study, we hypothesized that human labor onset is associated with changes in the abundance and posttranslational modification (e.g. phosphorylation) of key regulatory myometrial proteins. Specifically, we sought to characterize labor-associated changes in the hSHP αB-crystallin and HSP27 in human myometrium.

Materials and Methods

Experimental subjects

All experimental procedures performed in this study were approved by the University of Newcastle Ethics Committee in accordance with the institutional guidelines of the John Hunter Hospital, Australia, and the KK Women’s and Children's Hospital, Singapore. Informed consent was obtained from women before elective or emergency cesarean section. Myometrial tissue samples were crushed under liquid nitrogen and then homogenized in a urea/CHAPS-based lysis buffer (7 M urea, 2 M thiourea, 4% wt/vol 30 mM CHAPS, pH 8.5) at a ratio of 100 mg crushed tissue/1 ml extraction buffer. The homogenate was centrifuged (13,000 x g for 15 min at 4 C), and the resulting supernatant was collected. Depletion of HSA and IgG was performed using the albumin and IgG removal kit (GE Healthcare, Piscataway, NJ) according to the manufacturer's protocol. Briefly, extracts were incubated with an anti-HSA/anti-IgG resin for 30 min at room temperature. Unbound proteins were then separated from the absorption matrix using a mini-spin cartridge and centrifuging the sample at 1000 x g for 2 min. Concentration and desalting of HSA/IgG-depleted samples was performed by acetone precipitation. Four volumes of ice-cold acetone were added to each sample, incubated for 3 h at −20 C, and then centrifuged (13,000 x g for 15 min at 4 C). Samples were resuspended in 50 μl extraction buffer. Protein concentration estimation was performed using the 2-D Quant Kit (GE Healthcare, Uppsala, Sweden).

Extraction of myometrial proteins and depletion of human serum albumin (HSA) and IgG

Myometrial tissue samples were crushed under liquid nitrogen and then homogenized in a urea/CHAPS-based lysis buffer (7 M urea, 2 M thiourea, 4% wt/vol 30 mM CHAPS, pH 8.5) at a ratio of 100 mg crushed tissue/1 ml extraction buffer. The homogenate was centrifuged (13,000 x g for 15 min at 4 C), and the resulting supernatant was collected. Depletion of HSA and IgG was performed using the albumin and IgG removal kit (GE Healthcare, Piscataway, NJ) according to the manufacturer’s protocol. Briefly, extracts were incubated with an anti-HSA/anti-IgG resin for 30 min at room temperature. Unbound proteins were then separated from the absorption matrix using a mini-spin cartridge and centrifuging the sample at 1000 x g for 2 min. Concentration and desalting of HSA/IgG-depleted samples was performed by acetone precipitation. Four volumes of ice-cold acetone were added to each sample, incubated for 3 h at −20 C, and then centrifuged (13,000 x g for 15 min at 4 C). Samples were resuspended in 50 μl extraction buffer. Protein concentration estimation was performed using the 2-D Quant Kit (GE Healthcare, Uppsala, Sweden).

Fluorescent cyanine (Cy)-dye labeling of samples

Before labeling, Cy dyes Cy2, -3, and -5 (GE Healthcare) were reconstituted in anhydrous dimethylformamide. Nonlaboring (n = 6) and laboring (n = 6) myometrial proteins (50 μg) were labeled with 400 pM of either Cy3 or Cy5 dye on ice for 30 min in the dark. To account for any dye binding bias, half of every sample group was labeled with Cy3, whereas the remaining half was labeled with Cy5. An internal pooled label was also prepared. Myometrial proteins were then labeled with Cy2, Cy3, and Cy5 in a 1:1:1 ratio. The labeled proteins were mixed with the Cy2-labeled pooled control and made up to 450 μl with rehydration buffer (7 M urea, 2 M thiourea, 4% wt/vol CHAPS, 2 mg/ml dithiothreitol, and 1% vol/vol pH 3–10 immobilized pH gradient buffer).

Two-dimensional difference gel electrophoresis (2D-DIGE)

In-gel rehydration of immobilized pH gradient strips (nonlinear, pH 3–10, 240 mm; GE Healthcare) were performed with the Cy-labeled samples overnight for 12 h at room temperature under mineral oil (Sigma Chemical Co., St. Louis, MO). Isoelectric focusing was performed using a Multiphor II unit (GE Healthcare) for a total of 58,800 Vh using the following parameters: hold at 300 V for 900 Vh, ramp and hold at 1000 V for 3900 Vh, ramp and hold to 8000 V for 13,000 Vh, and hold at 8000 V for 41,000 Vh. Strips were then equilibrated and reduced in equilibration buffer (30% vol/vol glycerol, 2% wt/vol SDS, 7 M urea, trace bromophenol blue) with 0.5% dithiothreitol for 15 min at room temperature and then in equilibration buffer containing 4.5% iodoacetamide for 15 min at room temperature. Strips were then placed on homogenous 10% acrylamide gels (255 × 205 mm) cast in low-fluorescence plates using the Ettan DALTSix Electrophoresis Casting System (GE Healthcare). This produced six simultaneously cast gels using the same batch of acrylamide gel stock solution for each gel. Second-dimensional electrophoretic separation was carried out at 2.5 W/gel constant power for 30 min and then 100 W (total) constant power using a peltier-cooled Ettan DALT II electrophoresis unit (GE Healthcare) until the bromophenol dye front reached the bottom of the gel.

Imaging and analysis of 2D-DIGE gels

Fluorescence image acquisition was performed using the Typhoon 9400 Variable Mode Imager (GE Healthcare). Each gel was scanned at the following excitation/emission wavelengths: 480/530 nm (Cy2), 520/590 nm (Cy3), and 620/850 nm (Cy5). Gel analysis was conducted using the DeCyder version 5.0 Biological Variation software module (GE Healthcare) for spot matching, quantitation, and standardization of protein spot data. Spot matching was validated by manual inspection of gel spots. Statistically significant protein abundance changes were identified via Student’s t test. Proteins present in either all gels or at least in all nonlaboring or laboring gels and whose abundance changed more than 1.5-fold (P < 0.05) were chosen as proteins for identification.

Matrix-assisted laser desorption / ionization time-of-flight (MALDI-ToF) protein sequencing

A preparative sequencing gel was prepared by performing 2D-PAGE on 500 μg of pooled myometrium proteins. The gel was fixed in 50% methanol/7% acetic acid for 30 min and then stained with SYPRO Ruby (Invitrogen, Carlsbad, CA) overnight at room temperature. The gel was then washed in 10% methanol/7% acetic acid, briefly rinsed in distilled water, and visualized using a UV transilluminator. Target proteins were excised and destained in 50 mM ammonium bicarbonate in 50% vol/vol methanol for three washes of 40 min each. Gel plugs were dried at 37 C for 3 h and then incubated in 40 ng/ul trypsin (Promega, Madison, WI) in 20 mM ammonium bicarbonate at 37 C for 3 h. Sequencing was performed using an Ettn MALDI-ToF Pro (GE Healthcare). For this, 1 μl trypptic peptide was mixed with 1 μl α-cyano-4-hydroxycinnamic acid matrix (5 mg/ml in 50% acetonitrile, 0.1% trifluoroacetic acid), and 1 μl of this mixture was spotted in duplicate onto the MALDI target tray. Peptide mass fingerprinting was performed alongside peptide standards in reflectron mode. Sequence data were used to interrogate the Swiss-Prot and NCBInr databases. As an additional control measure, the isoelectric point and molecular weight of identified proteins were checked against the region of the pick gel from which they were excised.

One-dimensional (1D) SDS-PAGE and immunoblotting

Myometrial proteins (10 μg) were loaded on precast 12% NuPAGE acrylamide gels (Invitrogen) and separated electrophoretically using the Xcell SureLock Mini-Cell system at 190 V (constant) until the bromophenol blue dye migrated to the bottom of the gel. Proteins were then transferred to nitrocellulose (Hybond C; GE Healthcare) using the Xcell II blot module (Invitrogen) at 30 V for 55 min. Membranes were blocked with 5% skim milk in Tris-buffered saline supplemented with 0.01% Tween 20 (TBST) for 2 h and then incubated with antibodies for αB-crystallin (1:3000) Stresses (Stressgen Bioreagents, Victoria, Canada), total HSP27 (1:2500), pHSP27-Ser15 (1:1000), p-HSP27-Ser78 (1:2000), and p-HSP27-Ser82 (1:2000) in 5% skim milk/TBST for 2 h at room temperature. HSP20

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and HSP27 antibodies were purchased from Upstate Biotechnology (Lake Placid, NY). Membranes were then washed five times (5 min per wash) in TBST, and the appropriate secondary antibody (1:2000) was added for 1 h at room temperature. Immunoreactive protein was detected with enhanced chemiluminescence (GE Healthcare) using the Intelligent Dark Box LAS-3000 Imager (Fuji Photo Film, Tokyo, Japan). Membranes were stripped with 0.2 m NaOH and then reprobed with α-smooth muscle actin (α-SMA, 1:2000; Sigma) as a loading control. α-SMA abundance of actin protein in human myometrium remains unchanged with and during pregnancy (22, 23). Similarly, in rat myometrium, α-SMA mRNA and protein levels remain unchanged throughout gestation and labor onset (24). HSP27 phosphorylation was also standardized to α-SMA because its relative molecular mass (M₉ = 42 × 10⁶) differs from that of HSP27 and its phospho-forms (M₉ = 27 × 10⁶). Thus, we were able to avoid any issues of inhibited interaction between the antibody and the ligand during reprobing caused by residual HSP27 phospho-antibody binding. Specificity of immunoreactive bands was assessed by probing with preimmune antiserum followed by secondary antibody and also secondary antibody alone. Densitometric analysis was performed using Multi-Gauge (Fuji Photo Film) image analysis software. Band intensity values of α-b-crystallin and HSP27 were standardized to those of α-SMA. Statistical analysis was performed with GraphPad Instat version 3.0 (GraphPad Software, San Diego, CA).

Coimmunoprecipitation

Myometrial proteins (~100 mg, n = 5 for both nonlaboring and laboring samples) were extracted in 1 ml Nonidet P-40 (NP40) lysis buffer (150 mm NaCl, 50 mm Tris-HCl, 5 mm EDTA, 0.5% NP40). One microgram of rabbit-anti-HSP27 antibody (Upstate Biotechnology) was added to 50 μg extracted protein. As a control, 1 μg normal rabbit IgG was added to 50 μg pooled nonlaboring and laboring myometrial protein extracts. The protein-antibody mixture was incubated overnight at 4°C, and then 20 μl protein A-agarose immunoprecipitation reagent was added and incubated for 1 h at 4°C. The mixture was centrifuged at 1000 × g for 5 min at 4°C, and the resulting pellet was washed three times for 5 min each with 1 ml NP40 lysis buffer. The pellet was resuspended in 60 μl 1× lithium dodecyl sulfate buffer (Invitrogen) and heated at 70°C for 10 min. Samples were then centrifuged at 12,000 × g for 10 min and the supernatant collected. Immunoprecipitated protein (20 μl) was resolved by 1D SDS-PAGE, transferred to nitrocellulose, and probed with antibodies for α-b-crystallin, HSP27, and α-SMA.

Immunofluorescence microscopy and colocalization analysis

Myometrial tissue obtained at cesarean section (n = 6 for nonlaboring and laboring) were immediately fixed in neutral-buffered formalin for 48 h and then embedded in paraffin. Sections of 10 μm were attached to low-fluorescence glass slides. Slides were deparaffinized by incubating in the following solutions for 5 min each: 2 × 100% xylene, 3 × 100% absolute alcohol, 80% ethanol, and 70% ethanol. Antigen retrieval was performed by microwave heating the slides in citrate buffer (pH 6) at 95°C. Sections were blocked in 10% fetal calf serum in PBS for 30 min and then incubated in α-b-crystallin (1:400), HSP27 (1:400) and α-SMA (Sigma, 1:500) antibodies diluted in 10% fetal calf serum. Alexa Fluor 488 goat antimouse IgG and Alexa Fluor 594 goat antirabbit IgG (Invitrogen; both 1:200) were used as secondary antibodies. 4′,6-Diamidino-2-phenylindole (300 nm) was used as a nuclear stain. As controls, sections were incubated with preimmune sera and/or secondary antibodies alone. Only specimens demonstrating low background with control staining were used for analysis. Slides were mounted for imaging by adding 15 μl Vectashield H-1000 (Invitrogen) mounting media. Immunofluorescence microscopy was performed using a Zeiss AxioCam MRm(v2) and AxioPlan 2 epifluorescent microscope (Carl Zeiss, Sydney, Australia). Specimens were visualized using band pass filter sets that demonstrate negligible spectral overlap. Image registration in each channel was calibrated using a Focalcheck fluorescence microscope test slide (no. 1; Molecular Probes, Invitrogen). For each patient pair, nonsaturated images were collected with identical microscope and camera settings using the Axiovision software package (version 4.4). Colocalization analyses were then performed using the Colocalization Module of the software that creates a scatterplot of each channel. Defining a threshold creates a colocalization mask to represent high-intensity green and red pixels. The same threshold values were used for each sample pair comparison.

Isometric tension recordings

Myometrial samples were dissected into strips (7 × 2 × 2 mm), connected to a Grass FT03C force transducer (Grass Instruments, Quincy, MA), and suspended in organ baths containing 15 ml Krebs’ physiological salt solution. Strips were maintained at 37°C and pH 7.4 and continuously gassed with 95% O₂/5% CO₂. Passive tension of 1 mN was applied to each strip, and strips were allowed to equilibrate for 60–90 min until the development of spontaneous regular contractions. Strips were then exposed to a single dose of oxytocin 10⁻⁷ m for 5–20 min (until maximal tension reached) and immediately frozen in liquid nitrogen. Contractility was measured by integrating the area under the tension curve, using a Makal 8E data acquisition system with Chart software (version 5; ADI, Melbourne, Australia). Responses to oxytocin were compared as a percentage of spontaneous activity before treatment. Total and phosphorylated HSP27 was assessed using immunoblotting as previously described.

Results

Protein profiles of term nonlaboring human myometria (n = 6) and term spontaneously laboring myometria (n = 6) were quantitatively compared using 2D-DIGE (25) (Fig. 1A). One of the most prominent changes detected in protein abundance was a 69% decrease (Student’s t test, P < 0.05) in the sHSP α-b-crystallin (accession no. P02511) in laboring myometria (Fig. 1B and Table 1). Consistent with these results, immunoblotting for α-b-crystallin in a larger sample cohort showed a 71% decrease (Mann-Whitney U test, P < 0.01) in laboring myometria (n = 11) compared with nonlaboring tissue (n = 13) (Fig. 1D). The 2D-DIGE study also identified three isoforms of HSP27 (accession no. P04792) resolved in the form of a charge-train, indicative of posttranslational modification (Fig. 1C and Table 1).

To determine whether α-b-crystallin and HSP27 interact as reported for other tissues (7, 10), coimmunoprecipitation was employed. HSP27 immunoprecipitates from nonlaboring (n = 5) and laboring myometria (n = 5) were resolved by 1D SDS-PAGE, transferred to nitrocellulose, and probed with antibodies for HSP27 and α-b-crystallin (Fig. 2A). A single band of approximate relative molecular mass (M₉) of 23 × 10³ corresponding to α-b-crystallin was detected in nonlaboring and, to a lesser extent, laboring myometria. In contrast, no band was detected in the negative control (preimmune rabbit IgG), suggesting specific association between α-b-crystallin and HSP27 in these samples. Because HSP27 is a known actin-binding protein that may mediate smooth muscle contraction through stabilization of the actin filaments, we also probed the HSP27 immunoprecipitates for α-SMA (Fig. 2A). A positive immunoreaction was observed in both nonlaboring and laboring myometria, indicating specific interactions between HSP27 and actin in the human myometrium at the time of labor.

Coimmunofluorescence was then performed to examine the cellular localization of α-b-crystallin, HSP27, and α-SMA in the myometrium at labor. In nonlaboring myometria, α-b-crystallin was primarily localized to the perinuclear region (Fig. 2B). A similar pattern of staining was observed for HSP27. By applying a colocalization algorithm that identifies regions of containing, high levels of colocalization between α-b-crystallin and HSP27 were demonstrated in the both the perinuclear and cytoplasmic regions of nonlaboring myocytes. This result is consistent with...
the coimmunoprecipitation of these two proteins. This relationship was completely different in laboring samples where HSP27 was detected in a fibrillar arrangement throughout the perinuclear and cytoplasmic regions, which was highly colocalized with α-B-crystallin (Fig. 2C). In these tissues, α-B-crystallin was barely detectable above background levels.

Phosphorylation of HSP27 has been shown to promote the dissociation of sHSP oligomeric complexes (7, 9) and modulate the ability of HSP27 to interact with actin (11, 18). We hypothesized that changes in the relocation of HSP27 to the actin cytoskeleton in the myometrium during labor are associated with changes in its phosphorylation status. Immunoblotting with phosphospecific antibodies (HSP27-Ser15, -78, and -82) revealed marked changes in HSP27 phosphorylation at labor relative to α-SMA (Fig. 3A). Although total abundance of HSP27 was unchanged, HSP27-Ser15 was 3.0-fold higher in laboring myometria (Mann-Whitney U test, P < 0.05; Fig. 3B). In contrast, levels of HSP27-Ser82 were 85% less in laboring myometria (Mann-Whitney U test, P < 0.01). There was no significant change in HSP27-Ser78.

To confirm that the changes in HSP27 were related to contractile events, phosphorylation of HSP27 was also measured in vitro after treatment of nonlaboring myometrial strips with the uterotonic oxytocin (Fig. 4). Oxytocin treatment showed an average 206 ± 64% (mean ± sem, n = 5) increase in contractility above baseline. This was associated with a significant increase in HSP27-Ser15 phosphorylation compared with matched controls (paired Student’s t test, P < 0.05; Fig. 4, B and C).
phorylation of HSP27-Ser78 and -Ser82 did not alter with oxytocin treatment.

**Discussion**

We describe for the first time in the human myometrium labor-associated changes in the sHSP αB-crystallin and HSP27. We provide evidence that changes in these proteins are associated with actin cytoskeletal remodeling at the time of labor and use in vitro stimulation of myometrial muscle strips to show these changes are directly related to contraction.

As revealed using 2D-DIGE, one of the most significant changes in protein abundance in the human myometrium at labor is a 69% decrease in αB-crystallin (Fig. 1A). Corroborative evidence was provided using immunoblotting in a larger sample population (Fig. 1B). Scant information exists regarding the function and regulation of αB-crystallin in smooth muscle; however, in skeletal muscle, αB-crystallin acts as a chaperone protein providing cells with cytoprotection during stress by preventing protein aggregation (26). Evidence suggests αB-crystallin may also be important in the regulation of intermediate filaments (27–29). Mutations in the αB-crystallin gene lead to the breakdown and aggregation of intermediate filaments and the development of skeletal and cardiomyopathies (30–32). Decreased levels of αB-crystallin in actively contracting myometria are not consistent with such observations and suggest the protein may have a different function in myometrial smooth muscle.

Specific interactions between αB-crystallin and HSP27 have previously been reported in muscle cells (33). We found that
these proteins interact primarily in the cytoplasm of nonlaboring myometria (Fig. 2, A and B). Most likely due to decreased levels of αB-crystallin, this interaction is almost completely lost with labor (Fig. 2C). The functional significance of αB-crystallin/HSP27 complexes in human myometria is unclear; how-ever, we propose that decreased expression of αB-crystallin at the time of labor liberates HSP27 enabling it to participate in other cellular events such as cytoskeletal remodeling.

HSP27 can modulate cytoskeletal dynamics through the F-actin stabilization and microfilament organization (11, 29, 34). Phosphorylation of HSP27 in human platelets increases F-actin formation (35), whereas HSP27 phosphorylation is required for growth factor stimulation of F-actin formation in cultured cells.

**Fig. 3.** Immunoblot analysis of total and phosphorylated forms of HSP27 in nonlaboring (NL) and laboring (L) myometria. A, Representative immunoblots demonstrating the abundance of total and phosphorylated forms of HSP27 (pHSP27-Ser15, pHSP27-Ser78, and pHSP27-Ser82) in nonlaboring (NL, n = 13) and laboring (L, n = 11) myometrium. B, Densitometric analysis of band intensity revealed that pHSP27-Ser15 was 3-fold higher (*, P < 0.05, Mann-Whitney U test) in laboring myometria. In contrast, pHSP27-Ser82 abundance was decreased 84.62% (**, P < 0.01, Mann-Whitney U test) in laboring samples. Abundance of pHSP27-Ser78 remained unchanged with labor. Quantitation of HSP27 immunoreactive bands was performed by standardizing to the band intensity of α-SMA. Mr, Relative molecular mass x 1000; M, molecular weight marker.

**Fig. 4.** Changes in phosphorylation of HSP27 in oxytocin-stimulated contractions in nonlaboring myometrial strips. A, Representative tracing of spontaneous and oxytocin-stimulated contractions in nonlaboring myometrium. There was an increase in contractility of 206 ± 64% (mean ± SEM, n = 5) above baseline values. B, Immunoblotting of pHSP27-Ser15 abundance in control (C) and matched oxytocin-treated (T) myometrial muscle strips. C, Densitometric analysis of the immunoblots revealed a significant 2.7-fold increase (P < 0.05) in pHSP27-Ser15 after oxytocin treatment. Band intensity of pHSP27-Ser15 was standardized to the band intensity of α-SMA. Mr, Relative molecular mass x 1000; M, Molecular weight marker.
(15, 21). This suggests HSP27-actin interaction is modulated by phosphorylation. Consistent with a role in actin remodeling, we found that HSP27 is relocated from the cytoplasm of nonlaboring myometria, where it is associated with αβ-crystallin, to the actin cytoskeleton in laboring myometria (Fig. 2, B and C). Moreover, we observed that this relocation occurs simultaneously with increased HSP27-Ser15 phosphorylation and concurrent HSP27-Ser82 dephosphorylation at labor. Similar results have been reported by in the rat myometrium where HSP27-Ser15 phosphorylation is increased at the end of pregnancy (36). HSP27-Ser15 phosphorylation is thought to facilitate binding to and subsequent stabilization of actin filaments (18), a notion consistent with increased HSP27-Ser15 phosphorylation at labor. The significance of HSP27-Ser82 dephosphorylation in the myometrium is unknown but it is plausible that such a modification would confer a conformational change upon the protein, potentially modulating its interaction with other sHSP and/or components of the actin cytoskeleton.

In vitro treatment of nonlaboring myometrial muscle strips with the uterotonin oxytocin increased contractility 206% above baseline (Fig. 4A). These samples exhibited significantly higher levels of HSP27-Ser15 phosphorylation, confirming the association between HSP27-Ser15 phosphorylation and myometrial contraction. It has previously been reported that stimulation of rat aortic vascular muscle contraction with angiotensin II induces HSP27 phosphorylation (37). Likewise, the vasconstrictor throrbin increases HSP27 phosphorylation in both mouse cardiac myocytes (38) and aortic smooth muscle cells (39). Typically, these agonists have been shown to induce HSP27 phosphorylation via signal transduction cascades involving either p38 MAPK and its downstream targets MAPK-activated protein kinase-2 and -3 (37, 40, 41) or through protein kinase C activation (42). Consistent with a proposed role in smooth muscle contraction, the inhibition of HSP27 phosphorylation in rat aorta smooth muscle cells inhibits angiotensin II-induced contraction (37). Therefore, the evidence supports a critical role for HSP27 phosphorylation in myometrial contraction.

Our results support the hypothesis that smooth muscle contraction is more than the regulation of intracellular Ca2+ concentrations and the regulation of MLCK activity by second messenger systems. Our findings suggest that modulation of actin cytoskeletal assembly by sHSP may be an important additional regulated step. This new knowledge could have important consequences as the pathways regulating the phosphorylation and synthesis of sHSP are uncovered in different tissues. In the myometrium, it seems that an old uterotonin oxytocin may have a new modus operandi as a regulator of HSP27 phosphorylation.

Acknowledgments

We thank T. Engel, T. Finnegan, and the staff of the Obstetrics and Gynaecology Divisions at the John Hunter Hospital, Newcastle, Australia, and KK Children's and Women's Hospital, Singapore, for patient recruitment and sample collection. We also acknowledge R. F. Thorne from the Cancer Research Unit at the University of Newcastle, New South Wales, Australia, for materials and advice regarding immunofluorescence.

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Disclosure Statement: The authors of this manuscript have nothing to declare.

Received May 21, 2007. Accepted September 25, 2007.

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