# Expression of Nrf2 and its downstream gene targets in hibernating 13-lined ground squirrels, Spermophilus tridecemlineatus

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Abstract Mammalian hibernation is associated with wide important for conformational stability of the protein at near variation in heart rate, blood ßow, and oxygen delivery to0 C body temperatures in the torpid state. tissues and is used as a model of natural ischemia/reperfu-

sion. In non-hibernators, ischemia/reperfusion is typicallyKeywords Oxidative stress Antioxidant defense associated with oxidative stress but hibernators seem ttschemia resistanceTorporDarousal cycle deal with potential oxidative damage by enhancing anti-NF-E2-related factor-2 Superoxide dismutase oxidant defenses in an anticipatory manner. The present eme oxygenaseAßatoxin aldehyde reductase study assesses the role of the Nrf2 transcription factor in leart

the regulation of antioxidant defenses during hibernation.

Nrf2 mRNA and protein expression were enhanced in selected organs of 13-lined ground squirr@permophilus Introduction

tridecemlineatusduring hibernation. Furthermore, Nrf2

protein in heart was elevated by 1.4D1.5 fold at multiplen order to survive winter, many small mammals use stages over a torporDarousal bout including during entryhibernation. The winter hibernation season consists of long long term torpor, and early arousal. Levels returned toperiods of deep torpor (lasting days to weeks) interspersed euthermic values when squirrels were fully aroused in in-by brief interbout periods (often 12D24 h) when animals terbout. Protein levels of selected downstream target generowarm to 37\( \text{D}3\) During torpor, metabolic rate is prounder Nrf2 control were also measured via immunoblotting oundly depressed, frequently to only 1D5% of the normal over the torporDarousal cycle in heart. Cu/Zn superoxideuthermic rate, and core body temperature (Tb) decreases dismutase and aßatoxin aldehyde reductase levels near ambient (often falling to 0£6). By hibernating, increased signiPcantly during entry into torpor and thersmall mammals can often save up\*to90% of the energy gradually declined falling to control levels or below in fully that they would otherwise need to remain euthermic over aroused animals. Heme oxygenase-1 also showed the saither winter months 1]. Transitions to and from the torpid trend. This suggests a role for Nrf2 in regulating the anti-state are closely regulated by strong reversible controls oxidant defenses needed for hibernation success. Hefart (e.g., protein phosphorylation or dephosphorylation) on the was amplified by PCR and sequenced. The deduced aminates of multiple energy-expensive metabolic processes as acid sequence showed high identity with the sequence from ell as by the enhanced expression of selected genes other mammals but with selected unique substitutions (e.gwhose protein products address specibe needs of the animal proline residues at positions 111 and 230) that might be the hypometabolic, hypothermic state of torpar \$1.

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Hibernating mammals show a strong reduction of heart rate and blood ßow during torpor. These conditions would be considered severely ischemic for non-hibernating species. However, studies with Arctic ground squirrels found no evidence of hypoxia during torpor itself but measurement of blood oxygen content indicated global hypoxia



conditions occurred during the arousal period caused by increased expression of downstream gene products that are demand for oxygen that outstripped delivery capacity [ under Nrf2 control such as Cu/Zn SOD and HO-1. The Thus, the torporĐarousal cycle has features of an ischemiaresent study of 13-lined ground squirre permophilus reperfusion cycle which is known to cause oxidative stress ridecemlineatus substantiates this hypothesis. Nrf2 gene in other systems (e.g., human heart attack or stroke). Indeed nd protein expression were elevated during torpor and this because of this hibernating mammals are being used as correlated with a signibcant increase in the expression of stroke model system [. Another factor that can contribute downstream antioxidant enzymes that are known targets of to the generation of reactive oxygen species (ROS) in the large.

hibernator is the altered composition of lipid reserves that is

necessary to maintain lipid ßuidity at low Tb values. For

optimal hibernation, lipid depots must contain elevatedMaterials and methods

levels of polyunsaturated fatty acids (PUFAs) such as lin-

oleic acid [5] but PUFAs are very susceptible to free radical Animals

attack leading to autoxidation and the generation of lipid

peroxide radicals 1. Hence, because of these factors that Thirteen-lined ground squirrels (130-180 g) were captured increase susceptibility to ROS damage during the hiberby a licensed trapper (TLS Research, Michigan) and nating season, mammalian hibernators must set up efficient ansported to the Animal Hibernation Facility (NIH, antioxidant defenses to deal with wide variations in ROS Bethesda, MD). Hibernation experiments were conducted generation over torpor Darousal cycles. by the laboratory of Dr. J. M. Hallenbeck (National Insti-

Regulation of antioxidant defenses is under the controtute of Neurological Disorders and Stroke). Animals were of speciPc transcription factors that regulate gene expreshept on a fall day/night light cycle in shoebox cages sion, one of the prominent signal transduction pathwaysnaintained at 21C and fed ad libitum until they entered being the Nrf2/ARE pathway. The NF-E2-related factor-2 and Pnished the pre-hibernation phase of hyperphagia that (Nrf2) is a basic leucine zipper transcription factor that maximizes lipid stores. To was measured with an binds to a cytoplasmic repressor protein, Keap1, or tdmplantable Programmable Temperature Transponder different nuclear binding partners. When bound to Keap1PTT-200 (Bio Medic Data Systems). In order to implant and sequestered in the cytoplasm, Nrf2 is unable to stimthe temperature transmitters, ground squirrels were anesulate transcription. Nrf2 can also be directed towardshetized with 5% isoßuorane and then the transmitter was degradation by the proteasome via its interaction withinjected subcutaneously into the middle of the back using a Keap1 Bl. However, under oxidative stress conditions, sterile disposable syring ₱ ₽. When squirrels had reached Nrf2 is released from Keap1 and translocates into the plateau weight gain of 220D240 g, they were randomly nucleus 91 where it dimerizes with proteins such as the divided into groups for one of two protocols. Initial trials to small Mafs, Jun and activating transcription factor-4 (ATF-characterize organ-specibe expressionnoff2 compared 4) [10D12]. The complex then binds to the antioxidant two groups. Euthermic controls were maintained under the response element (ARE) in the promoter region of genesame conditions as previously and Tb was conbrmed as that respond to oxidative stress. Well-known Nrf2-36D38C at the time of sampling. The other group was responsive antioxidant genes include thioredoxin and GSTslaced in a dark chamber at @. Most animals entered [13, 14], Cu/Zn superoxide dismutase (Cu/Zn SOD) deep torpor within 3D8 days and were sampled after each and heme oxygenase-1 (HO-1) 6. Furthermore, the individual had been hibernating for 2D5 days (as indicated enzyme aßatoxin aldehyde reductase (AFAR1) is an alddey continuous Tb readings of 6 C). Euthermic control keto reductase that is involved in the detoxipcation ofanimals were sampled on the same days as the hibernators. various aldehydes and ketones including the environmentall animals were anesthetized and then sacribced by carcinogen aßatoxin £117. A recent study has shown that decapitation. Tissues were excised, frozen immediately in the enzyme also contains multiple AREs in its promoterliquid nitrogen and then transported to Ottawa on dry ice region [18]. where they were stored at 80 C until use.

The potentially harmful oxidative stress conditions that In subsequent studies, pre-hibernation treatment was the hibernators may experience over the course of torpor ame but animals were sampled at multiple time points arousal cycles led us to hypothesize that the Nrf2 pathwayover the course of torpor and arousal in &C4dark cold well-known to be involved in ROS detoxiPcation in room. In this protocol, all animals were transferred into the mammals, would play a role in providing the hibernator cold room at the start of the experimental course. The with the necessary antioxidant defenses required to dealxperimental groups sampled were: (1) active in the cold with hibernation-related oxidative stress. This couldroom (ACR); these euthermic squirrels had not yet entered include an increase in Nrf2 levels during hibernation and torpor after 3 days in the cold as determined by Tb



readings, (2) early entrance into torpor with falling Tb with 294 ml of DEPC treated water. Ethidium bromide (ENT) (Tb = 31D12C, >14 h in the active state after full (0.3 mg/300 ml) was added in the solution and the mixture arousal from a torpor phase of at least 5 days), (3) early invas heated. The heated solution was then poured in a gel the torpid state <24 h) with a stable Tb of 5Đ7C (HIB-E), (4) later in the torpid state (at least 3 d continuousaliquot of PCR product was mixed with 2 of 6x blue/ torpor) with stable Tb of\* 5D7C (HIB-L), (5) early in arousal with rising Tb (AR-E) (Tb= 9D12C), and (6) fully aroused in interbout with Tb back at 37 for at least 18 h before sampling (AR). Animals in the aroused groupsbromide on a UV box. The bands from the most dilute had been in torpor continuously for at least 3 d.

Total RNA isolation and quality assessment

Total RNA was isolated from tissue samples using TrizolServices (Ottawa, ON). The sequence was conbrmed as reagent (Gibco BRL), according to manufacturer Osncoding Nrf2 by sequence comparison in BLAST. instructions, and resuspended in diethylpyrocarbonate (DEPC) treated water. RNA concentration was determined Vestern blotting by absorbance at 260 nm and the ratio of absorbance at 260/280 nm was used as an indicator of RNA purity. RNAFrozen tissue samples (500 mg) were homogenized in quality was also assessed by running samples on a 1.22ml of buffer containing 100 mM MOPS, 25 mM HEPES, agarose denaturing gel and staining with ethidium bromid 25 mM  $\beta$ -glycerophosphate, 5 mM EDTA, 1 mM EGTA, to reveal two sharp bands in every sample.

cDNA synthesis and PCR amplibcation of Nrf2

to amplify both Nrf2 andα-tubulin. The primers used for Storey [20] with 10% gels (5% stacking gel), 2μ0g of Nrf2. The forward primer sequence was TSCCCAGGT TGCCCACAT-3 and the reverse was ATGCCRGAG TCAGARTC-39. As a control genex-tubulin was amplified with forward (59-AAGGAAGATGCTGCCAATAA-39) and reverse (\$GGTCACATTTCACCATCTG-3) primers. The PCR was performed by mixing \$\mathbf{f}\$ of each cDNA dilution with 1.25 µl of primer mixture (0.5µM forward and 0.5  $\mu M$  reverse), 15  $\mu l$  of sterile water, 2.5  $\mu l$  of 10  $\times$ PCR buffer (Invitrogen), 1.25 of 50 mM MgCl<sub>2</sub>, 0.5 µl of 10 mM dNTPs, and 0.125l of Tag Polymerase (Invitrogen) for a total volume of 251. The cycles performed for amplibcation consisted of an initial step of 2 min at 94 followed by 94C for 1 min, 54C for 1 min, and 72C for 1 min repeated 37 times; the Pnal step was atC72or 1× TAE buffer prepared by mixing 6 ml of 50 TAE

and 250µM Na<sub>3</sub>VO<sub>4</sub>, adjusted at pH 7.4, with 1 mM phenylmethylsulphonyl ßuoride added immediately before homogenization. After centrifugation at 10,000 of or 10 min at 4C, supernatants were collected and soluble A 30 µg aliquot of total RNA from heart of hibernating protein concentrations were determined using the Coosquirrels was used for Prst strand cDNA synthesis usingnassie blue dye-binding method and the BioRad prepared Superscript II reverse transcriptase (Invitrogen) and fol-reagent (BioRad, Hercules, CA). SDS-polyacrylamide gel lowing the manufacturer Os protocol. Serial dilutions of the lectrophoresis and blotting to polyvinylidene dißuoride cDNA in water were prepared (10010-4) and were used membranes was carried out essentially as in Morin and amplibcation of Nrf2 were designed using the Primerprotein loaded per well, and electrophoresis at 200 V for Designer program, v3.0 (ScientiPc and Educational Soft45 min. Wet transfer of proteins onto PVDF membranes ware) based on the consensus sequences of mammalianas made using a transfer buffer solution containing 25 mM Tris (pH 8.5), 192 mM glycine and 10% v/v methanol at 4C for 1.5 h at 0.3 mA. Following transfer, membranes were blocked for 1 h in TBST (50 mM TrisĐ HCl pH 6.8, 150 mM NaCl, 0.05% v/v Tween 20) with 2.5% w/v powdered skim milk. This was decanted and then membranes were incubated overnight at 4with primary antibodies. Antibodies specibc for mammalian Nrf2 were purchased from Santa Cruz Biotechnologies and used at a 1:200 v:v dilution in TBST. Antibodies specibo to Cu/Zn SOD and HO-1 were purchased from Stressgen and used at dilutions of 1:4000 and 1:2000, respectively. AFAR1 antibody was a gift from Dr. John D. Hayes, University of Dundee and was used at a 1:1000 dilution. Subsequently, membranes were incubated with HRP-linked anti-rabbit 2 min. PCR products were separated on a 1.0% agarose glad G secondary antibody (1:2000 v:v dilution) in TBST for The gel was prepared by adding 3 g of agarose to 300 ml of h and then blots were developed using the SuperSignal West Pico Chemiluminescent Substrate (Pierce) according buffer (242 g Tris base, 57.1 ml concentrated acetic acidto the manufacturerÖs protocol. Bands were visualized 100 ml of 0.5 M EDTA in 1 I water, adjusted to pH 8.5) using a ChemiGenius Bio Imaging System (SynGene, MD,

tray and the gel was allowed to cool and solidify. A 110

on the 1% agarose gel. The gel was run in TAE buffer.

orange loading dye (Promega) and the solution was loaded

After separation, the bands were visualized with ethidium

cDNA sample were used for quantiPcation purposes to make sure that the products had not reached amplibication

saturation. Anrf2 fragment of 717 bp was found and was excised and sequenced by Canadian Molecular Research

USA) and band densities were quantiped using the assaccession number DQ328859. Figureshows the partial ciated Gene Tools software.

#### Quantibcation and statistics

Bands corresponding torf2 PCR product were normalized against bands for-tubulin amplibed from the same cDNA against a group of three Coomassie stained protein band that did not show variation between the different experi-DunnettÖs test for the hibernation cycle data.

Results

## cDNA cloning of Nrf2

translated amino acid sequence of ground squirrel Nrf2 aligned with the sequences for the human, mouse and rat protein. The full Nrf2 sequence has 605 residues in humans and 597 residues in mice and rats whereas the amplibed portion of S. tridecemlineatus Nrf2 encoded 239 amino acids residues, corresponding to 40% of the full sample. Western blot band intensities were standardized sequence. Ground squirrel Nrf2 was quite similar to Nrf2 fom other mammals and shared 87.9, 77.4, and 78.3% identity to human, mouse and rat Nrf2, respectively, over mental states and were not located close to the protein. Ground squirrel Nrf2 contained a few bands of interest. SigniPcant differences between the... groups were assessed using GraphPad InStat 3 software Fig. 1. These included substitutions of two proline residues ibernating mammals; these are shown in bold underline in hibernating data or analysis of variance followed by the at positions 111 and 230 as compared with the human sequence. An asparagine residue replaced a tyrosine residue at position 226 and a tyrosine was substituted for a histidine at position 227.

Nrf2 gene expression

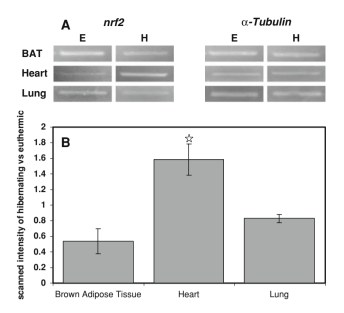
Levels of nrf2 mRNA transcripts were measured for three tissues of euthermic and hibernating ground squirrels:

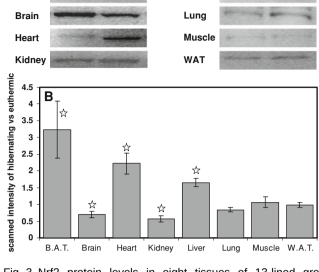
Using RT-PCR and primers derived from the consensus rown adipose tissue (BAT), heart and lung. Using RT-PCR sequence of frf2 from other mammalian species, a PCR and the primers for ff2 and alpha-tubulin, a constitutively product of 717 bp was retrieved from total RNA prepared expressed gene, the relative levels of both transcripts were from heart of hibernating ground squirrels. The productassessed in each organ. Levels not 2 transcripts were was conprimed as encoding a portion of three sequence normalized against the tubulin transcript level in each and the sequence was submitted to GenBank withsample. Figureb shows the ratio of normalized of 12

Fig. 1 Partial amino acid sequence (239 residues) of Nrf2 from 13-lined ground squirrels (S. tridecemlineatuscompared to Nrf2 sequences of human (Homo sapien)s mouse Mus musculus and rat Rattus norvegicus; Genbank accession numbers for the four species are DQ328859, NP\_006155, NP\_035032 and O54968 respectively. The full length protein is 605 amino acids for human Nrf2 and 597 amino acids for mouse and rat Nrf2. Dashed lines in the alignment represent amino acids that are identical with the ground squirrel sequence; spacer dots are inserted when residues are not present in all sequences. Selected substitutions in the. tridecemlineatussequence are shown in bold underline

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squirrel
                                   mmdlelpppglpsqqdmdlidilwrqdidlgvsrevfdfsqrrkeyelekqkklekerqeqlqkeqekaffaqlqldeet 80
human
                                   \verb|mmdlel|| pppg|| qsqqdmdlidi|| wrqdidl|| gvsrevfdfsqrqkdyelekqkklekerqeq|| qkeqekaffaqfqldeet 80 | wrddidliger 
                                   mmdlelpppglqsqqdmdlidilwrqdidlqvsrevfdfsqrqkdyelekqkklekerqeqlqkeqekaffaqlqldeet 80
                                                        ......SQVAHIPKPDALYFDDCMQLLAETFPFVDDNEVSSATFQSLVPDIPSHIESPVFNAPP 58
squirrel
human
                                   geflpiqpaqhiqtdtsgsasy-----q----e----
                                                                                                                                                                                          --h-....--al---
mouse
                                   geflpiqpaqhiqtdtsgsvsy-----
squirrel
                                   OAOSPETSLDGAMA.DLNNIOODIEOVWOELFSIPELOCLNIENDKLVETTTVPSPEAKLTEID.NNYFYPSIPSLEKEV 136
human
                                                     ---vaqvapv--dgm----e--l----
mouse
                                   ----lns--ea-t.--ss-e--m-----t--kq-ad--a----t--m-s-yh--s--s----- 232
                                   ----lds--et--t.--ss---m------t--kqqa------t--m-s-yh--s------ 232
rat.
                                   GNCSPHFLNAFEDSFSSILSTEDPNQLTVNSLNSDATLNTDFGDEFYSAFIAEPSTSNSMPSSATVSQSLSELLYG.
 squirrel
                                                                                                                                           human
                                    ---g----hg------d-as--..t--d-np--
mouse
                                    ds-----hg-----d-as--..--d-np------1----ggg-
squirrel
                                   SDLSLCKAFNONHPESTAEFNDSDSGI..................
                                   -----slntspsvaspehsvesssygdtllglsdseveeldsapgsvkqngpktpv.h 398 c----pk-a-g-m----slntspsvaspehsvessiygdpppgfsdsemeeldsapgsvkqngpkaqpah 390
mouse
                                   \verb|c------| k-t-g-v-------s| lntspsraspehsvessiygdpppgfsdsemeeldsapgsvkqngpkaqpth 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 390 | 3
rat
squirrel
                                   ssqdmvqplspsqqqsthvhdaqcentpekelpvspqhrktpftkdkhssrleahltrdelrakalhipfpvekiinlpv 478
human
                                   spgdtvqplspaqghsapmresqcenttkkevpvspghqkapftkdkhssrleahltrdelrakalhipfpvekiinlpv 470
mouse
                                    ssgdtvqplspaqghsaavhesqcenttkkevpvspghqkvpftkdkhssrleahltrdelrakalhipfpvekiinlpv 470
squirrel
                                    vdfnemmskeqfneaqlalirdirrrgknkvaaqncrkrkleniveleqdldhlkdekekllkekgendkslhllkkqls 558
mouse
                                   ddfnemmskegfneaglalirdirrrgknkvaagncrkrklenivelegdlghlkderekllrekgendrnlhllkrrls 550
                                   ddfnemmskegfneaglalirdirrrgknkvaagncrkrklenivelegdlghlkderekllrekgendrnlhllkrkls 550
rat
squirrel
                                    tlylevfsmlrdedgkpyspseyslggtrdgnvflvpkskkpdvkkn 605
human
mouse
                                     tlylevfsmlrdedgkpyspseyslqqtrdgnvflvpkskkpdtkkn
                                    tlylevfsmlrdedgkpyspseyslqqtrdgnvflvpkskkpdtkkn 597
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Liver

Fig. 2 Effect of hibernation omrf2 mRNA levels in three tissues of Fig. 3 Nrf2 protein levels in eight tissues of 13-lined ground 13-lined ground squirrels.a) Representative bands on agarose gelssquirrels. a) Representative Western blots show expression of the Paired tubes ampliÞendrf2 and α-tubulin transcripts from the same samples andnrf2 band intensities were normalized against the squirrels; animal conditions are as in Fig. 2. Data are measEM, tissue. b) Histogram showing the ratio of normalized PCR product ing euthermic controlP < 0.05 levels in tissues from hibernating (2Đ5 days of continuous torpor in the 4 C cold room with Tb readings of 6 C) versus euthermic (at 21 C with Tb 36Đ38C) ground squirrels. Values are meahsSEM for n=3 independent trials on tissue from different animals nation cycle: (1) euthermic squirrels that were active in the \*, Hibernating values are signibcantly different from the correspond-cold room (ACR) and had not yet entered hibernation, (2) ing euthermic controlP < 0.05

show PCR product levels amplified from ground squirrel cDNA. Nrf2 band at 57 kDa. ti) Histogram shows mean relative protein expression in tissues from hibernating versus euthermic ground  $corresponding \hbox{$z$-tubulin intensity for the same sample. Band sizes } n=3 \quad independent \quad trials \quad on \quad tissue \quad from \quad different \quad animals.$ were 717 bp fonrf2 and 616 bp fox-tubulin. BAT is brown adipose \*, Hibernating values are signiPcantly different from the correspond-

levels at six different points over the course of the hiber-

early entrance into torpor with falling Tb (ENT), (3) early in the torpid state <24 h) with a stable Tb at 5Đ7C

(HIB-E), (4) later in the stable torpid state (at least 3 d)

rising Tb (AR-E), and (6) fully aroused in interbout with Tb back at 37C (AR) (Fig. 4). Nrf2 protein in heart of

ground squirrels entering hibernation rose signipcantly to

transcript levels in the tissues. Relative levels of 2 transcripts changed significantly only in heart, with a 1.6-foldwith Tb at \* 5Đ7C (HIB-L), (5) early in arousal with increase in transcript levels during hibernation.

### Nrf2 protein expression

levels that were 1.36-fold higher than the values in ACR Nrf2 protein levels were measured by immunoblotting.controls P < 0.05). Amounts remained elevated through-Western blot analysis of Nrf2 in ground squirrel tissuesout torpor and early arousal (by 1.3 to 1.5-fold compared revealed that the content of the 57 kDa Nrf2 proteinwith ACR) but declined back to control levels in fully increased signibcantly in BAT, heart and liver during aroused animals. hibernation (by 3.2-, 2.2- and 1.6-fold, respectively, In order to determine whether elevated Nrf2 during

RΔT

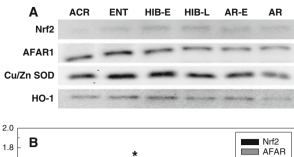
and 60% of the euthermic values, respectively < 0.05).

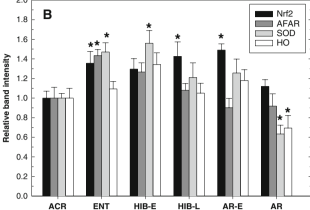
Protein levels of Nrf2 and downstream targets in hibernating heart

The above data for bothmrf2 mRNA and Nrf2 protein euthermic values. Cu/Zn SOD levels also rose during levels indicated that changes in this transcription factor arentrance into hibernation by 1.5-foldP (< 0.05) and important in heart during hibernation. In order to further remained high during hibernation. Levels decreased sigexplore Nrf2 expression in heart, we measured Nrf2 proteimibcantly when squirrels were fully aroused from torpor;

P < 0.05) (Fig.3). Interestingly, Nrf2 protein levels hibernation altered the expression of genes that are under decreased in brain and kidney of hibernating animals to 70Nrf2 control in other mammals, levels of three proteins were assessed in heart over the same time course of hibernation (Fig.4). AFAR1 protein levels in ground squirrel heart increased signipcantly during entrance into hibernation by 1.43-fold as compared with ACR animals (P < 0.05). Levels then declined gradually back to







Nrf2, AFAR1, Cu/Zn SOD and HO-1 in other mammalsb) ( (fully debned in the Materials and Methods): active in the cold room importance of this transcription factor in protecting cells (ACR), entrance into hibernation (ENT), early-hibernation (Hib-E), the corresponding values for ACR. Data are mean  $\mathbf{SEM}$ , n = 3D7independent trials on heart samples from different animals. \*, Value is strain [30]. Furthermore, hyperoxia-induced levels of signibcantly different from the corresponding ACR valæç 0.05

heart SOD levels in the AR group were just 63% of the Nrf2 knockout mice than in control mice, linking Nrf2 to ACR value P < 0.05). HO-1 protein showed an increasing the antioxidant defenses needed to deal with hyperoxiatrend during hibernation (although not significant) but, likeassociated ROS productions 1. Hence, Nrf2 and its mals (AR), falling to 69% of the ACR value P(< 0.05).

## Discussion

ground squirrels 11. Studies with black bears have also highlighted increases in a marker for lipid peroxidation during hibernation [22]. Indirect evidence that oxidative stress occurs comes from several studies of antioxidant defense mechanisms which have shown specibc elevation during hibernation of the activities of selected antioxidant enzymes including glutathione peroxidase in ground squirrels Citellus citellus [23], peroxiredoxin in both bats (Myotis lucifugus and ground squirrelsS). tridecemlineatus) [24, 25], and serum catalase and superoxide dismutaselike activity in hamsters Mesocricetus auratus 26, 27, Furthermore, plasma ascorbate levels in Arctic ground squirrels Spermophilus parry)i rose by 3Đ5 fold during each torpor bout and were then rapidly depleted during arousal, the highest rate of decrease correlating with the time when oxygen consumption was maximal during thermogenesis28. Microanalysis of tissue and extracellular Buid in brain of hamsters also showed depletion of ascorbate during arousal, consistent with oxidative stress in this stage of the hibernation cyclegi.

Clearly, then, hibernators need well-developed antioxi-Fig. 4 Protein expression of Nrf2, AFAR1, Cu/Zn SOD and HO-1 in dant defenses and this would predictably include increased heart of S. tridecemlineatus (a) Representative Western blots expression of genes coding for antioxidant enzymes. One showing single strong cross-reacting bands at 57, 40, 19 and way to stimulate the antioxidant response is through activation of transcription factors that regulate these genes. Histogram showing relative protein expression in squirrels held in aThe present study examines the expression of the tran-4 C cold room and sampled from different physiological conditions scription factor Nrf2. Many studies have shown the late-hibernation (Hib-L), early-arousal (Ar-E), and fully aroused against oxidative stress. For example, studies using Nrf2 (AR). Data for standardized band intensities are expressed relative tonockout mice showed that expression of several detoxi-Þcation enzymes was strongly reduced in the knockout mRNA for antioxidant enzymes such as NAD(P)H:quinone oxidoreductase 1 (NQO1) and HO-1 were much lower in

SOD, HO-1 decreased signibcantly in fully aroused ani-downstream enzymes could be good indicators of current or anticipated oxidative stress in hibernator tissues. The Nrf2 transcription factor from ground squirrel heart

was partially cloned in the N-terminal region of the protein. The amplibed fragment was 239 amino acids long and started at position 103 as compared with the human Nrf2

The potential for oxidative stress during the different sequence. Several amino acids substitutions were identibed hibernation stages has been assessed previously. Most when the ground squirrel partial Nrf2 sequence was comthe work performed to date has focused on the levels and/orared to Nrf2 sequences from non-hibernating mammals. the activity of antioxidant enzymes and markers of anti-The amplibed Nrf2 segment from ground squirrels conoxidative defense over the hibernation time course. Thisained two key domains, Neh4 (within residues 98D156) study adds to this previous work by measuring the levels oand Neh5 (within residues 153D227), which have both been the Nrf2 transcription factor and of its target genes at dif-shown to act synergistically to allow Nrf2 transactivation ferent time points over the torpor Darousal cycle. Evidence through CREB binding 32. Deletion of these domains that oxidative stress occurs during hibernation has beeleads to a severe disruption in Nrf2 transactivation activity. produced previously from studies with intestine of 13-linedIt has also been demonstrated that phosphorylation of



residues within these domains by mitogen-activated kinasectivated under many conditions that lead to ROS generfamily members, ERK and/or JNK, leads to a positive ation such as anoxia, hypoxia, and ischema, [38]. Control of HO-1 involves transcription factors including regulation of Nrf2 transactivational activity38. Only minor modifications were seen in the Neh4 and Neh5HIF-1 and Nrf2 [16, 38]. Cu/Zn SOD is a cytoplasmic domains of the ground squirrel protein suggesting that thenzyme that catalyzes the breakdown of harmful superoxtransactivation process is largely conserved in the hiberide into hydrogen peroxide and water. It is involved in the nator. However, two changes of interest occurred nearegulation of ROS generation and the Cu/Zn SOD gene is these regions: the substitutions of proline residues for app-regulated in response to oxidative stress. Cu/Zn SOD serine/arginine residue at position 111 and for a serine an also help to reduce cellular injury following reperfuresidue at position 230. These modibcations, along with theion of tissues that were previously ischemical since other substitutions that were not exclusive to the ground vrf2 was elevated over the hibernation time course, protein squirrel, could result in a conformational change to thelevels of these downstream enzymes were also measured hibernator Nrf2 protein that might aid its transcriptional over the same time course to see if they correlated with activity at low body temperatures. Nrf2 levels. The data showed that Cu/Zn SOD and AFAR1

Levels of nrf2 mRNA transcripts were measured in three protein levels increased significantly in parallel with eleground squirrel tissues using RT-PCR. Transcript levelsyated Nrf2 levels during entry into torpor whereas HO-1 were signibcantly elevated only in heart during hibernatiorprotein levels also showed an upward trend (Alg.This (by 1.6-fold) whereas amounts in other tissues appeared touggests that induction of synthesis of these antioxidant be down-regulated. Previous studies have been contradiproteins is triggered as one of the Prst events when animals tory as to whethen rf2 transcript levels rise under oxidative begin to suppress metabolic rate. This highlights the posstress conditions. Whereas Cho et al.1][ reported an sibility that hibernators increase antioxidant defenses as an induction of nrf2 transcripts by 2.0- and 2.6-fold in lungs of anticipatory or preparatory response in order to deal with mice subjected to 48 and 72 h of hyperoxia, a study bythe oxidative stress that occurs during prolonged torpor and Papaiahgari et al.341 had contrary results. Data gathered arousal. Such a preparation for oxidative stress by elevation here seems to agree with the former study single gene of antioxidant defenses has been proposed previously for induction was observed. Nrf2 protein levels were thenother situations of hypometabolism/arousal such as during measured in ground squirrel tissues by immunoblottingestivation in land snails40. It is interesting that none of Levels were signipcantly elevated in BAT, heart and liverthe three downstream proteins remained signipcantly ele-(3.2-, 2.2- and 1.6-fold, respectively) during hibernation vated in heart during early arousal (although Nrf2 was), a but decreased significantly in ground squirrel brain and ime when oxidative stress should be high due to the high kidney. rates of oxygen-based thermogenesis needed to return to

Due to the concomitant increase of Nrf2 mRNA and euthermic Tb [28, 29]. Hence, it is possible that these protein levels in ground squirrel heart during hibernation, particular Nrf2 regulated enzymes actually have their most Nrf2 protein levels in this organ were assessed at multiple prominent functional role during the torpor period itself. time points over a hibernation cycle. SigniPcantly elevated/arious defensive mechanisms are needed to ensure metlevels of this transcription factor were found over hetero-abolic stability and viability over the long term during thermic portions of the time course: entry into torpor, latetorpor, since the potential to degrade oxidatively damaged torpor, and arousing from torpor (Fig.). Levels returned proteins and/or synthesize new proteins is strongly supto near control values only after animals were fully pressed during torpor as part of the general suppression of aroused. Reports have suggested that Nrf2 protein levelsTP expenditure [3]. However, given that Nrf2 levels increase under oxidative stress[36] 36]. Hence, the general remain elevated during early arousal, it is probable that elevation of Nrf2 protein over the torpor portions of the additional gene targets under Nrf2 control may remain hibernation cycle suggests that the antioxidant defenses effected and/or be speciPcally up-regulated during the hibernator heart may also be enhanced during this time. arousal phase. One of these may be peroxidredoxin (Prdx).

In order to assess this idea, three enzymes that have recent report documented Nrf2 control over Prdx1 in ARE binding sites in their promoter regions were choserresponse to hypoxia/reoxygenation1. Our analysis of for study: Cu/Zn SOD, AFAR1 and HO-1. These enzymesPrdx isozymes in 13-lined ground squirrels showed strong all play important roles in the detoxiPcation of ROS (SOD,hibernation responsive up-regulation of 3 Prdx isozymes in HO-1) or xenobiotic carbonyl compounds (AFAR1) and heart including a 13-fold increase in Prdx1 protein over they could all have roles in the protection against oxidative euthermic levels [15]. Nrf2 may be the transcription factor stress in hibernators. HO-1 is an inducible enzyme that is esponsible.

involved in heme degradation and the subsequent produc- Overall, these results document an up-regulation of the tion of biliverdin, a known antioxidant [7]. The enzyme is Nrf2 transcription factor in heart and selected other organs



during hibernation suggesting that gene targets under Nrf23. Kim IC, Masutani H, Yamaguchi Y, Itoh K, Yamamoto M, Yodoi control have important roles to play in hibernation success. This is substantiated by parallel increases during hibernation in Cu/Zn SOD and AFAR1 protein levels, as well as peroxiredoxins 25, all known Nrf2 gene targets. We 14. Chanas SA, Jiang Q, McMahon M, McWalter GK, McLellan LI, conclude that a Nrf2-mediated up-regulation of multiple antioxidant enzymes is a key preventative defense against oxidative stress occurring over the torporDarousal cycle.

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