Osteopathy and resistance to vitamin D toxicity in mice null for vitamin D binding protein

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A line of mice deficient in vitamin D binding protein (DBP) was generated by targeted mutagenesis to establish a model for analysis of DBP’s biological functions in vitamin D metabolism and action. On vitamin D–replete diets, DBP+/− mice had low levels of total serum vitamin D metabolites but were otherwise normal. When maintained on vitamin D–deficient diets for a brief period, the DBP+/−, but not DBP+/+, mice developed secondary hyperparathyroidism and the accompanying bone changes associated with vitamin D deficiency. DBP markedly prolonged the serum half-life of 25(OH)D and less dramatically prolonged the half-life of vitamin D by slowing its hepatic uptake and increasing the efficiency of its conversion to 25(OH)D in the liver. After an overload of vitamin D, DBP−/− mice were unexpectedly less susceptible to hypercalcemia and its toxic effects. Peak steady-state mRNA levels of the vitamin D–dependent calbindin-D9K gene were induced by 1,25(OH)2D more rapidly in the DBP−/− mice. Thus, the role of DBP is to maintain stable serum stores of vitamin D metabolites and modulate the rates of its bioavailability, activation, and end-organ responsiveness. These properties may have evolved to stabilize and maintain serum levels of vitamin D in environments with variable vitamin D availability.


Introduction
Vitamin D binding protein (DBP), also known as the group-specific component of serum (Gc-globulin), is a member of the albumin, α-fetoprotein, and α-albumin multigene family (1, 2). DBP is a highly polymorphic serum protein predominantly synthesized in the liver as a single-chain glycoprotein of ~58 kDa (3). The serum concentration of human DBP is 4–8 μM and its serum half-life is 2.5–3 days. DBP has a high-affinity binding site for 25(OH)D3 (5 × 10^−8 M^−1), the major circulating form of vitamin D that is generated by 25-hydroxylation of vitamin D in the liver. This site also binds 1,25(OH)2D3, the active form of the vitamin, as well as the parent vitamin D itself, both with somewhat lower affinity (4 × 10^−8 M^−1) (4). Vitamin D sterols are necessary to maintain normal serum calcium homeostasis and bone development. Deficiency of vitamin D results in the bone diseases of osteomalacia and rickets, diseases characterized by formation of poorly calcified and structurally impaired bones. DBP has several biological activities in addition to its ability to bind vitamin D. DBP binds avidly to G-actin (2 × 10^−8 M^−1) via a binding domain in its carboxy terminus (5, 6); this binding can sequester circulating monomeric G-actin, preventing polymerization into F-actin after cellular trauma (7). DBP can activate macrophages in vitro (8) and enhances C5a-mediated chemotaxis (9). A definitive approach to testing the multiple function(s) of DBP in vivo, and most particularly its role in vitamin D metabolism and action, has not been available.

The role of DBP in vitamin D action is not defined. However, it is likely that it functions in accordance with the “free-hormone” hypothesis. This hypothesis suggests that plasma sterols are trapped in the vascular compartment via their association with a corresponding set of serum binding proteins. These complexes provide an available reservoir of steroid hormones; the sterols can be made available to cells by their dissociation from the binding protein (10). Additional roles of the binding proteins might include facilitation of sterol/steroid entry into the cell, as suggested by the identification of cell surface receptors for certain serum steroid binding proteins (11, 12); facilitation of its intracellular bioactivity and gene activation; or protection from excess free hormone (13).

The highly polymorphic structure of DBP has led to its use as a serum protein marker for population-genetics studies. Sera from many thousands of individuals from all five continents have been studied for variations in the electrophoretic mobility of DBP (14). Despite this intensive analysis, no individuals have been identified who are null for DBP. This observation has led to the suggestion that one or more functions of DBP may be essential to human viability (4). In this report, a mouse line lacking DBP was successfully established by targeted disruption of the endogenous murine DBP gene.
These mice have been used to test essential role(s) of DBP and specifically to delineate the role(s) played by DBP in vitamin D metabolism and action.

Methods

Materials

All restriction and modification enzymes were purchased from New England Biolabs (Beverly, Massachusetts, USA). Chemicals and tissue culture supplies were purchased from Sigma Chemical Co. (St. Louis, Missouri, USA) unless otherwise stated. CDNA probe labeling kits were purchased from Boehringer-Mannheim Biochemicals (Indianapolis, Indiana, USA). Oligonucleotides were synthesized by the Nucleic Acid Core Facility of the Children's Hospital of Philadelphia (Philadelphia, Pennsylvania, USA). [3H]vitamin D3 (15 Ci/mmol) and 25(OH)[26,(27)-methyl-3]vitamin D3 [25(OH)[3H]D3; 15 Ci/mmol] were purchased from Amersham Life Sciences Inc. (Arlington Heights, Illinois, USA). The [3H]vitamin D3 used at the beginning of the study went out of production during the course of these experiments, and studies were completed using [3H]vitamin D3 (3.85 Ci/mmol) provided by Michael Holick (Boston University, Boston, Massachusetts, USA). 1,25(OH)2 vitamin D3 [1,25(OH)2D3] was provided by Milan Uskokovic (Hoffman-La Roche, Nutley, New Jersey, USA). Hybridization probes in these studies were as follows: an 865-bp rat DBP EcoRI cDNA fragment (2), a 180-bp mouse calbindin-D28k EcoRI cDNA fragment (clone B6; gift from Elizabeth Bruns, University of Virginia, Charlottesville, Virginia, USA), and a 400-bp mouse-rL32 EcoRI– HindIII fragment (15).

Generation of DBP–/– mice. The mouse (m) DBP gene was cloned from a 129 Sv mouse genomic library (Lambda FXII vector; Stratagene, La Jolla, California, USA), using the 875-bp 5' end of the rat (r) DBP cDNA as hybridization probe. Four clones were obtained. Exon content and organization were determined by restriction mapping and hybridization to exon fragments of the homologous rDBP gene (16). An 8.8-kb EcoRI fragment of the 24-kb “clone 2” containing exons 2–8 was selected for generation of the targeting vector. A 1.8-kb SalI-XhoI fragment containing the PGK-promoter/neomycin phosphotransferase (PGK-neo5) cassette was blunt-end ligated into exon 5 of the mDBP fragment at the BamHI site to generate pDBPN5. To shorten the construct, pDBPN5 was digested with SalI and KpnI by removing a 2.3-kb fragment containing exons 2 and 3 generating pDBPN5(4–8). The diphtheria toxin A chain gene (DTA) cassette (17) (generous gift from Barbara Knowles, The Jackson Laboratory, Bar Harbor, Maine, USA) on a 2.7-kb SalI-KpnI fragment was ligated to the KpnI-SalI pDBPN5(4–8) fragment, and circularized recombinants were selected. This procedure resulted in the targeting vector, pDBPN5(4–8)-DTA.

The targeting vector was transfected into passage 9 R1 embryonic stem (ES) cells (a gift from Andras Nagy, Mount Sinai Hospital, Samuel Lunenfield Research Institute, Toronto, Ontario, Canada). The ES cells were grown on mitomycin C–treated SL2 feeder cells plated on gelatin-coated plates (18). The ES cells were maintained in the differentiated state by growth in LIFD cell (Genetics Institute, Cambridge, Massachusetts) conditioned media. R1 cells (0.8 ml; passage 13) at a concentration of 7 × 104/ml were electroporated with 40 µg of linearized pDBPN5-DTA using a Gene Pulser (Bio-Rad Laboratories, Hercules, California, USA) at 240 V/500 µF and replated onto feeder cells. G418 (GIBCO BRL, Gaithersburg, Maryland) at 200 µg/ml was added 48 h later. After 8 days of culture, resistant colonies were picked and transferred to 96-well plates. After splitting into triplicate 96-well plates, DNA was isolated from lysed cells, digested with EcoRI, resolved on 0.8% agarose gels, and Southern blotted with a probe generated from mDBP exon 2.

One hundred ninety-five surviving ES clones were analyzed, and three had been successfully targeted. Clone D1 was expanded, and 10–15 cells were microinjected into 3.5-day-old blastocysts from a C57BL/6J mouse by the University of Pennsylvania Transgenic and Chimeric Mouse Core. Five chimeric mice resulted, and four transmitted the targeted mDBP gene to their agouti offsprings after backcross to C57BL/6J mates. The mouse colony was maintained within a microbiological barrier facility, and animals were anesthetized with tribromoethanol (Avertin) at 300 mg/g body weight during all invasive procedures and before sacrifice.

Genotyping

PCR assay. Tail DNA (1 µg) was subjected to PCR with two sets of oligonucleotide primers. The first set included primers from exon 5, DBP-A (5'-CGCGCTCTGCACTTGTGGTG-3') and DBP-B (5'-GCATACTTTGGTTGTTGCAG-3'). This primer set spanned the neo' cloning site and generated a 100-bp fragment only from the DBP+/+ allele. A second, confirming primer set included DBP-C, also from exon 5 (5'-CTCTGGCACTTTAGTGCTTAC-3'), and DBP-D, derived from neo' gene sequences (5'-GGATGGAGATGTTGTTGGA-3'). These primers generated a 120-bp fragment specific to the DBP–/– allele. For both sets of oligonucleotides, PCR was carried out over 30 cycles: 94°C for 1 min, 54°C for 30 s, and 72°C for 2 min.

Radial immunodiffusion assay. One percent agarose containing 3% polyclonal anti-rDBP antibody (cross-reactive with mDBP) was poured onto a glass backing, and circular wells were cut into the solidified matrix. Test mouse sera or controls (2 µl) were loaded into each well and allowed to diffuse for 48 h. The gels were rinsed for 24 h, first with PBS, and then with distilled water. Gels were stained with 0.2% Coomassie brilliant blue in 5% methanol and 5% trichloroacetic acid for 30 min, and then destained with 5% methanol and 7.5% acetic acid for 24 h and air dried. The diameters of the stained immunoprecipitated circles were proportionate to the amount of mDBP in each serum sample.

Western analysis. Mouse sera were resolved on 7.5% SDS-polyacrylamide gels. Gels were electrophoresed onto nitrocellulose membranes, incubated with rabbit anti-rDBP, and visualized by enhanced chemiluminescence (ECL kit; Amersham Life Sciences Inc.) as described previously (19).

25(OH)D binding assay. Mouse sera were analyzed for 25(OH)D binding by incubation with 25(OH)[3H]D3 in the presence of increasing concentrations of cold 25(OH)D3 as described previously (20). Scatchard analysis was carried out to estimate the mean binding capacity (21). Results shown were representative of three separate analyses, each carried out in duplicate.

RNA isolation and Northern analysis. RNA was extracted from primary tissues using TRIzol Ultrapure (Life Technologies Inc., Gaithersburg, Maryland, USA) according to the manufacturer's protocol. To obtain high-quality RNA from the intestine, the RNA extraction procedure was modified (22). RNA (10 µg) was denatured in 20% MOPS solution (0.04 M MOPS; 0.01 M sodium acetate; 2 mM EDTA, pH 7.0; 50% formamide; and 6.25% formaldehyde) at 65°C for 10 min. The RNA was separated in 1% agarose-formaldehyde gels, transferred overnight to ZetaBind membranes (CUNO Inc., Meriden, Connecticut, USA), and hybridized with 32P-labeled probes at 42°C for 16 h. After washing the filters, the radioactivity in each band was quantified by PhosphorImager (Molecular Dynamics, Sunnyvale, California, USA).

Blood chemistry. Mice were maintained for the noted time periods on either sterile standard laboratory diet or vitamin D–deficient diet (Purina test diet No. S826C-5: no vitamin D, calcium 0.60%, phosphorus 0.40%; North Penn Feeds Inc., Lansdale, Pennsylvania, USA). Animals were sacrificed and blood was collected by cardiac puncture. Plasma calcium, phosphorus, and alkaline phosphatase were measured using a Kodak Ektachem Analyzer. Parathyroid hormone (PTH) measurements by immune assay were kindly provided by the labo-
ratory of Leonard Deftos, University of California at San Diego, San Diego, California, USA (23). 1,25(OH)₂D₃ and 25(OH)D were measured by RIA (24, 25).

Bone histomorphometric analyses. Age- (6–8 months) and sex-matched DBP⁺⁺ and DBP⁻⁻ mice were maintained on standard laboratory chow (controls) or the vitamin D-deficient diet for 8 weeks in temperature-controlled rooms with 12-h light and 12-h dark cycles. Animals were fluorochrome double-labeled with calcine in neutral PBS (0.5 mg/animal) delivered intraperitoneally 10 days and 3 days before sacrifice. Femurs were removed and dehydrated in 70% ethanol for at least 3 days. Bone samples were embedded in methylmethacrylate, and 5-μm sections were stained with Masson’s trichrome (26) as described previously (27). They were examined using a semiautomated image analysis system (BIOQUANT IV; R&M Biometrics, Nashville, Tennessee, USA) by an investigator blinded to group allocations. Unstained sections (8–10 μm) were examined for fluorochrome-based indices. Histomorphometric analyses were performed using the Statview statistical package (Abacus Concepts Inc., Berkeley, California, USA) and presented in the format of Parfitt et al. (28).

Preparation of vitamin D sterols for intravenous injection. [³H]vitamin D₃ and 25(OH)[³H]D₃ were dried under a stream of nitrogen and stored in the dark at 4°C. These [³H]sterols were preincubated, at 4°C overnight ([³H]vitamin D₃) or for 1 h (25(OH)[³H]D₃), with aliquots of sera collected from either DBP⁺⁺ or DBP⁻⁻ mice. In this way, the sterols could be solubilized in either DBP-free or normal serum without using organic solvents. Approximately equal cpm were delivered using equal aliquots, indicating equivalent recovery. Such preincubated DBP⁺⁺ sera were used for injections into DBP⁺⁺ mice, and preincubated DBP⁻⁻ sera were used for injections into DBP⁻⁻ mice. 1,25(OH)₂D₃, stored dry under argon and shielded from light, was solubilized in ethanol, dried under nitrogen, and similarly incubated with homologous sera from DBP⁺⁺ or DBP⁻⁻ mice at 4°C overnight, and was injected for the calbindin-D₉₅ gene induction experiments.

Vitamin D kinetics. Age-matched groups of DBP⁺⁺ and DBP⁻⁻ mice were fed vitamin D₃-deficient diets for 4–6 weeks. Animals were fasted overnight before the injections. Each animal was weighed at the beginning of the experiment. [³H]vitamin D₃ or 25(OH)[³H]D₃ (0.5 μCi, or 100,000 cpm) in homologous sera was injected intravenously into the tail vein. Animals were sacrificed at different time points over 24 h. Blood, and in some cases liver, and/or kidneys were collected. Heparinized blood was spun at 2,500 rpm to collect the plasma, and aliquots were counted in BCS scintillant (Amersham Life Sciences Inc.). An aliquot of each collected tissue was weighed and incubated with NSCII tissue solubilizer (Amersham Life Sciences Inc.) at 1ml/100 mg tissue overnight at 50°C. After complete solubilization, the samples were cooled and neutralized with glacial acetic acid to decrease chemiluminescence. Final tissue counts were obtained using Ready-Organic scintillation fluid (Beckman Coulter Inc., Fullerton, California, USA) in a refrigerated scintillation counter after chemiluminescence had decayed to a plateau.

For analysis of urinary excretion, 100,000 cpm of 25(OH)[³H]D₃ preincubated with homologous sera was injected intravenously into anesthetized DBP⁺⁺ and DBP⁻⁻ mice.

Figure 1
Targeted disruption of the mouse DBP locus. (a) A fragment of mouse genomic DNA containing exons 4–8 of the DBP gene was used to design the targeting vector. A PGK-promoter/neomycin phosphotransferase cassette was inserted at the BamHI (M) site in exon 5 to disrupt the mDBP gene and provide for positive selection. A DTA cassette was ligated to the 5′ HindIII site for selection against random integration. (b) Restriction enzyme mapping distinguished the intact from the disrupted DBP allele. A mouse DBP exon 2 probe, located outside of the targeting vector itself, hybridized to an 8.8-kb EcoRI (R) fragment from the native DBP allele and a 5.7-kb EcoRI fragment from the disrupted mDBP allele. Restriction sites are: S, SalI; H, HindIII; B, BgII; R, EcoRI; C, ClaI; M, BamHI. (c) The targeting vector (α) was transfected into ES cells, G418 selection was applied, and surviving clones were analyzed by Southern blotting. Analysis of 8 representative ES cell lines among the 65 examined is shown. All eight clones contained the native 8.8-kb mouse DBP EcoRI fragment, and one clone (D1) also contained the 5.7-kb fragment, indicative of successful homologous recombination. DBP, vitamin D binding protein; DTA, diphtheria toxin A; ES, embryonic stem.
increasing concentrations of cold 25(OH)D3 is shown. The data were further analyzed and DBP–/– mice hybridized with rat DBP cDNA is shown. The full-length 1.8-kb mDBP mRNA was detected in wild-type mice, present at diminished levels in heterozygous mice, and totally absent in mice homozygous for DBP gene activation. Balanced RNA loading was confirmed by the ethidium bromide staining of 18S rRNA (bottom). (b) Western analysis for serum DBP in 12 mice representing each of the three genotypes is shown. The antibody was a cross-reacting, polyclonal rabbit antiserum to rat DBP. The presence and relative levels of the 58-kDa DBP paralleled the mRNA levels in panel a. The antibody was a cross-reacting, polyclonal rabbit antiserum to rat DBP. The autoradiogram of the Northern blot analysis of total RNA from livers of DBP+/+, DBP+/–, and totally absent in mice homozygous for DBP gene activation. Balanced RNA loading was confirmed by the ethidium bromide staining of 18S rRNA (bottom). (c) Semiquantitative radial immunodiffusion analysis of sera from mice of all three genotypes using the polyclonal antiserum confirmed the absence of DBP in DBP–/– mice. The antibody was a cross-reacting, polyclonal rabbit antiserum to rat DBP. The autoradiogram of the Northern blot analysis of total RNA from livers of DBP+/+, DBP+/–, and totally absent in mice homozygous for DBP gene activation. Balanced RNA loading was confirmed by the ethidium bromide staining of 18S rRNA (bottom). (d) Serum saturation binding analysis using tracer 25(OH)[3H]D3 in the presence of 25(OH)D3 and 25(OH)[3H]D3 was performed (Statview statistics package), and data were considered significant when P < 0.05.

Results

Generation of DBP–/– mice. A fragment of the mDBP gene spanning exons 2–8 was isolated from a mouse 129 SV genomic library by cross-hybridization with a rat DBP cDNA probe (Fig. 1a). The 8.8-kb genomic insert was restriction-mapped and partially sequenced to confirm its identity and establish restriction sites for vector construction. A vector for homologous recombination was generated by cloning the PGK-neo cassette into the BamHI site into DBP exon 5 and ligating a DTA cassette into intron 3 at the 5¢ terminus of the targeting insert (Fig. 1a). Mouse ES cells were electroporated with the targeting insert, and surviving cells (i.e., those lacking the DTA cassette) were selected for resistance to G418. One hundred ninety-five surviving ES colonies were analyzed by Southern blotting (Fig. 1, b and c). A hybridization probe from exon 2 detected an 8.8-kb EcoRI fragment from the intact mouse DBP gene and a 5.7-kb EcoRI fragment from the disrupted locus. Of three ES cell clones with the disrupt-
ed DBP allele, a single colony, D1, was selected for subsequent use (Fig. 1c). D1 ES cells were expanded and microinjected into 3.5-day-old blastocytes from a C57Bl/6 mouse; five chimeric animals that ranged from 5% to almost 100% agouti coat color were obtained. The near-100% agouti chimeric animals produced only agouti offspring, 56% of which carried the DBP allele. DBP+/+ animals were intercrossed; of the initial 177 offspring, 25% were wild type (DBP+/+), 49% were heterozygous (DBP+/–), and 26% were homozygous (DBP–/–) for the disrupted locus. This distribution, consistent with normal Mendelian inheritance, suggested that DBP–/– mice were of normal viability. Intercrosses of DBP–/– progeny resulted in normal frequency and sizes of litters, indicating normal fertility and fecundity of the DBP–/– males and females. DBP–/– mice appeared grossly normal, had growth curves identical to wild-type littermates, and showed no abnormalities after necropsy and histologic examination of all major organs (data not shown).

Verification of DBP-null state. Loss of expression from the disrupted DBP allele was confirmed by analysis of mRNA and protein. Northern blots of total liver RNA from DBP+/+, DBP+/–, and DBP–/– mice were generated. The expected 1.8-kb DBP mRNA was detected in the DBP+/+ liver, present, but at reduced levels, in DBP+/– liver, and totally absent from DBP–/– liver (Fig. 2a). Western blots of sera from animals of each genotype confirmed the reduction of DBP in DBP+/– mice and the total absence in DBP–/– mice (Fig. 2b). Analysis of DBP content in sera by radial immunodiffusion readily distinguished between the levels in the wild-type and carrier mice and confirmed the absence of immunoreactive DBP in DBP–/– mouse serum (Fig. 2c). The simple radial immunodiffusion assay was subsequently used for genotyping. The profiles of total serum proteins as determined by sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) were otherwise identical in DBP+/+ and DBP+/– sera (not shown). Serum saturation binding analysis using tracer 25(OH)[3H]D3 was performed to detect any bioactive fragments of DBP that might be produced by the mutant locus. No binding activity was detectable in the DBP–/– serum. The affinity constants were essentially identical in wild-type and heterozygous sera (Kd = 1.0 × 10−9 M and 0.9 × 10−9 M) and comparable to that previously reported for hDBP (5 × 10−9 M). From the Scatchard plot, the maximum binding capacity of the DBP+/+ serum was estimated as 570 μg/l and for DBP+/– serum, 380 μg/l (67% of wild-type) (Fig. 2d). When sera were preincubated with 25(OH)[3H]D3 and subjected to a native gel electrophoresis (32), the DBP band was absent from DBP+/– sera and detected at reduced levels in DBP–/– sera compared with that of DBP+/+ animals (data not shown). These data demonstrated a complete absence of expression of immunologically or biologically detectable DBP in the DBP–/– mice. The data also suggested that heterozygotes have a minor upregulation of protein expression from the intact DBP allele.

DBP+/– mice are vitamin D-depleted. The consequences of the DBP-null mutation on vitamin D levels and calcium homeostasis were determined. Serum levels of calcium, phosphorous, alkaline phosphatase, PTH, 25(OH)D, and 1,25(OH)2D were initially measured in DBP+/+, DBP+/–, and DBP–/– littermates maintained on standard laboratory chow containing vitamin D. On this standard diet, DBP–/– mice were vitamin D–deficient (vitamin D–) (Fig. 3, a and b). DBP+/– mice had significantly lower total serum levels of 25(OH)D and 1,25(OH)2D than DBP+/+ mice (Fig. 3, a and b). Heterozygotes fell within the intermediate range in both assays. No significant differences in serum calcium, phosphorous, or alkaline phosphatase were observed between wild-type and DBP–/– littersates on a vitamin D–deplete diet (Table 1), and the levels of PTH were equivalent for these two groups (Fig. 3e). Therefore, the low total serum sterol levels in the DBP–/– mice coexisted in equilibrium with adequate intracellular concentrations of 1,25(OH)2D under steady-state conditions, based on
analysis of the PTH response.

The mice were next stressed by placing them on a vitamin D–deficient diet for four to six weeks. The effects of this diet on calcium and vitamin D homeostasis were compared between the DBP+/+ and wild-type littermates. After four weeks without exogenous vitamin D, the 25(OH)D3 and 1,25(OH)2D3 levels decreased in the wild-type mice to levels comparable to those of the DBP+/+ animals, the lowest detection limits of these assays (Fig. 3, d and e). At this point, both groups became hypophosphatemic and alkaline phosphatase levels increased slightly (Table 1). Serum calcium was maintained at the same level in the DBP+/+ and DBP–/– groups. Although no differences between the two groups were noted in serum PTH levels while on standard diet (Fig. 3c), there was a doubling in the mean PTH levels in the DBP–/– group on the vitamin D–deficient diet (P < 0.05; Fig. 3f). Thus, the DBP+/+ mice were more sensitive to dietary vitamin D deficiency than their normal littermates, selectively manifesting secondary hyperparathyroidism.

Bone histomorphometric analyses demonstrated an increased sensitivity of the bone in DBP–/– mice to vitamin D deficiency. Skeletal radiographs showed no discernible differences between DBP+/+ and DBP–/– mice (data not shown). Qualitative bone histological examination and quantitative histomorphometric analyses were carried out to detect bony abnormalities at a higher level of resolution. The bones of age- and sex-matched DBP+/+ and DBP–/– littermates were studied on standard diets and after eight weeks on vitamin D–deficient diets. The following end points were analyzed: femoral lengths, trabecular bone volume/tissue volume (BV/TV) in two planes, osteoid surface/bone surface (OS/BS), osteoclast surface/bone surface (OC/BS), osteoclast number/bone surface (OCN/BS), mineralizing surface/bone surface (MS/BS), mineral apposition rate (MAR), and osteoid thickness (OTH). On standard diets, there were no significant differences between the DBP+/+ and DBP–/– groups in any of these end points (data not shown), nor was there any distinguishable difference in osteoclast numbers.

On the vitamin D–deficient diet, BV/TV was significantly lower in both the DBP+/+ and DBP–/– groups compared with values on the standard diet (8% vs. 4%, P < 0.01 for DBP+/+ groups; 12% vs. 4%, P < 0.05 for DBP–/– groups). This shared decrease confirmed the effectiveness of the vitamin D–deficient diet in altering bone metabolism without regard to the DBP status.

The remaining end points demonstrated a markedly increased and selective sensitivity of the DBP–/– mice to bone changes caused by vitamin D deficiency. OS/BS of the DBP–/– group was significantly higher on the vitamin D–deficient diet (from 3.3% to 8.8%, P < 0.001, male animals only; data not shown), whereas the OS/BS in the DBP+/+ group was not influenced by diet. A comparison between the two groups after vitamin D deprivation showed the net result of this differential effect (Fig. 4c, P < 0.05). The accentuated thickening of the osteoid seams in the DBP–/– mice, reflecting osteoblast activity and mineralization, could be appreciated qualitatively in a visual comparison of the Masson’s trichrome–stained sections (Fig. 4, a and b; arrowhead). This selective effect in the DBP+/+ mice was confirmed quantitatively by measurement of the mean OTh (Fig. 4d; P < 0.001). A significant and selective difference between the DBP+/+ and DBP–/– mice was also noted in the lower MAR and MS/BS in the DBP–/– mice (Fig. 4, e and f; P < 0.05 and P < 0.01, respectively). When fed a vitamin D–replete diet,
neither DBP+/+ nor DBP–/– mice had skeletal abnormalities. However, when fed a vitamin D–deficient diet, the DBP–/– group demonstrated abnormally high osteoblastic activity, with undermineralization of the newly synthesized matrix compared with the matched DBP+/+ group. Thus, the DBP–/– mice demonstrated an increased sensitivity to dietary vitamin D deprivation by developing hypovitaminosis D osteopathy.

DBP increases the serum half-life of vitamin D and 25(OH)D. DBP is the main carrier of vitamin D in the serum and as such has been proposed to play a role in plasma clearance of 25(OH)D (33). To directly establish the role of DBP in the clearance and distribution of 25(OH)D, DBP+/+ and DBP–/– mice were subjected to four weeks of vitamin D deprivation sufficient to generate serum 25(OH)D deficiency in both groups (Fig. 3, d and e). This period was followed by intravenous injection of tracer amounts of 25(OH)[3H]D3 previously incubated with homologous serum (see Methods). Animals were sacrificed over short (40 minutes) and long (24 hours) time courses, and plasma was collected and counted. In DBP+/+ mice, 25(OH)[3H]D3 was gradually cleared from plasma with ~15% of the injected isotope still present after 24 hours (Fig. 5a). In contrast, the half-life of 25(OH)[3H]D3 in the plasma of the DBP–/– mice was markedly shortened, with a very low level detected in the circulation several minutes after injection (Fig. 5a and inset).

The fate of the rapidly cleared 25(OH)[3H]D3 from the plasma of the DBP–/– mice was investigated. Urine was collected from the DBP+/+ and DBP–/– animals for 24 hours after intravenous injection of the tracer 25(OH)[3H]D3. Threefold more isotope was excreted into the urine of the DBP–/– group than the wild-type controls (Fig. 5b). Total urine 25(OH)D content was specifically quantified by radioimmunoassay (RIA) as well. A 2.4-fold greater clearance was observed over a 24-hour period in the DBP–/– mice (DBP+/+, 149 pg/24 hours; DBP–/–, 361 pg/24 hours). Aliquots of plasma harvested at each time point of the short time course (Fig. 5a and inset) were extracted and fractionated by TLC. The percentage of cpm migrating in the 25(OH)D region (left) and the polar region (right) of the chromatograph for each time point were plotted.
Figure 6
Accelerated entry of serum [3H]vitamin D into the liver and its conversion to polar metabolites. [3H]vitamin D3 was preincubated with aliquots of DBP+/+ or DBP–/– serum and injected intravenously into mice in the respective groups. Plasma samples (a) and livers (b) were harvested at the indicated times after injection, and tritium counts were obtained. Data were expressed as a percentage of total cpm injected normalized to total plasma volume (P < 0.05 at 20 and 40 min; a) or per gram of liver (b), and represent the mean ± SEM from three independent experiments. (c) Aliquots of plasma from the 1-min time point in a were extracted and subjected to TLC. The position of a 25(OH)D standard was localized by ultraviolet visualization, and the percentage of total chromatographed cpm migrating in the 25(OH)D region was plotted. (d) The percentage of total chromatographed cpm migrating in the polar region was plotted. The data are the mean ± SEM of two to three independent experiments.

had been preincubated in homologous serum. These animals were sacrificed over a period of 40 minutes, plasma was collected, and livers were homogenized. The clearance of [3H]vitamin D3 from the DBP–/– plasma was substantially more rapid than from the DBP+/+ plasma (Fig. 6a). A reciprocal pattern of [3H]vitamin D3 uptake was seen in the livers. Between 10 and 20 minutes after injection, tritium counts were 14–15-fold higher in the livers of the DBP–/– animals than in the DBP+/+ controls (Fig. 6b). By 40 minutes after injection, the level of tritium in the DBP–/– livers fell to a value indistinguishable from the DBP+/+ controls. This pattern suggested that the accelerated transfer of [3H]vitamin D3 from the serum to the liver of the DBP–/– mice was followed by egress of isotope from the liver. This accelerated entry into the liver appeared to be an effect specific to this organ because there was less isotope detected in the kidney parenchyma of DBP–/– animals at all time points (not shown).

The accelerated rate of transfer of vitamin D3 from the serum to the liver in the DBP–/– mice might be paralleled by the hepatocyte-mediated 25-hydroxylation to 25(OH)D or conversion of the vitamin D to water-soluble inactive esters that could then be excreted (34). To determine which pathway was being used, aliquots of plasma from the one-minute time points of each [3H]vitamin D3 kinetic experiment were extracted and subjected to TLC. One minute after injection, a mean of 32% of the total plasma tritium migrated at the 25(OH)[3H]D position on thin-layer chromatographic analysis in the wild-type animals compared with 7.5% in the DBP–/– animals (Fig. 6c). At the same time point, almost three times more polar metabolites were detected in DBP–/– mice than in DBP+/+ mice (29% and 11%, respectively). Thus, the accelerated entry of vitamin D into the liver in the absence of DBP appeared to result in the shunting of a substantial proportion of the substrate into an inactivating pathway or pathways. The relative contributions of liver and other tissues to this pathway cannot be determined from the present data.

The toxic response to high-level vitamin D administration was decreased in the absence of DBP. The profile of serum clearance and hepatic modification of vitamin D seen in the DBP–/– mouse suggested that DBP was necessary for efficient hepatic 25-hydroxylation and subsequent activation. This observation would suggest that the biologic activity of administered vitamin D would be significantly blunted in DBP–/– mice. This result would contradict the prediction from the free-hormone hypothesis that the lack of serum DBP in the DBP–/– mouse would accentuate the effects of a given dose of vitamin D due to absence of its putative buffering action (13). To differentiate between these two predictions, the effect of a sublethal dose (1000 IU/g body weight) of vitamin D3 was compared between wild-type and DBP-deficient mice. There were no significant differences in weight loss between the two groups over a one-week period of observation after delivery of vitamin D (data not shown). At day 7, the animals were sacrificed, serum was collected, and kidneys were harvested. The large dose of vitamin D resulted in the expected increase in serum calcium levels in the DBP+/+ mice compared with control mice injected with vehicle only. Remarkably, the DBP–/– mice responded to the same dose of vitamin D with a significantly less pronounced increase in serum calcium (Fig. 7a); von Kossa staining of kidneys confirmed the greater effect of the vitamin D dose on serum calcium levels in the DBP+/+ mice by demonstrating correspondingly more significant accumulation of calcium deposits in the cortices of the DBP+/+ kidneys (Fig. 7, b–e). Thus, the lack of serum DBP resulted in relative protection of the DBP–/– mice from the toxicity of high-dose vitamin D administration.

1,25(OH)2D-dependent gene induction is more rapid in the absence of DBP. DBP–/– mice were more susceptible to vitamin D deficiency and were relatively resistant to vitamin D toxicity (Figs. 3 and 7). This observation could imply that the bioactivity of 1,25(OH)2D is in some way dependent on DBP. The importance of DBP to the delivery of biologically active 1,25(OH)2D to target tissues and the activation of gene expression was tested. Induction of calbindin-D9K mRNA in the proximal small intestine is
dependent on 1,25(OH)₂D. The steady-state levels of proximal intestinal calbindin-D₉K mRNA were determined in vitamin D–deficient DBP⁺/⁺ and DBP⁻/⁻ mice 24 hours after intravenous injection of 1,25(OH)₂D. Before injection (t = 0), proximal intestinal levels of calbindin-D₉K mRNA were identical (Fig. 8, a and b). Immediately after injection of 1,25(OH)₂D, there was a transient depression in normalized calbindin-D₉K mRNA levels in both groups. This period was followed by a progressive increase to a peak value and a subsequent fall toward baseline values. The overall maximal levels at the peak of induction were similar in both groups, although the timing was different. In the DBP⁺/⁺ animals, calbindin-D₉K mRNA levels reached the maximum response 12 hours after 1,25(OH)₂D injection. This kinetic response was in agreement with that previously reported in wild-type mice by others (35). In contrast, the calbindin-D₉K mRNA levels in the DBP⁻/⁻ mice reached a similar maximal response at eight hours (Fig. 8, a and b). These kinetic differences were statistically significant and reproducible over four experiments. Thus, DBP slowed the kinetics but not the overall level of calbindin-D₉K gene expression in response to induction by 1,25(OH)₂D.

Discussion

A mouse model carrying a disrupted and functionally inactivated DBP locus was generated to investigate the role of DBP in vitamin D metabolism. The DBP allele was disrupted by targeted homologous recombination within exon 5 of the mouse DBP gene (Fig. 1). Assays on the sera of the mice homozygous for the disrupted DBP allele failed to detect immunoreactive DBP fragments or 25(OH)₂[D3]-binding peptides (Fig. 2, b–d). The interruption of the endogenous mouse DBP locus was therefore felt to effectively eliminate expression of DBP and high-affinity serum binding of vitamin D and its metabolites.

Based on extensive population surveys, it was initially suspected that homozygosity for a DBP-null allele might result in embryonic or neonatal lethality (4). The generation of DBP⁻/⁻ mice at the expected mendelian frequency from intercrosses between DBP⁺/⁻ heterozygotes was therefore unexpected. These DBP⁻/⁻ mice were normal in size and appearance, and intercrosses between DBP⁻/⁻ mice resulted in litters of normal size with no evidence of impairment of fertility in either males or females. It was concluded that a total absence of serum DBP was fully consistent with normal viability and reproduction.

The DBP⁻/⁻ mice had significantly lower serum levels of 25(OH)D and 1,25(OH)₂D compared with DBP⁺/⁺ animals (Fig. 3). The free-hormone hypothesis states that the biological activity of a given hormone is related to its unbound (“free”) rather than protein-bound concentration in the plasma (10). Thus, by binding to sterol or steroid hormones and trapping them in the vascular compartment, each binding protein creates a buffer or reservoir for its respective hormone (36). Normally, only 0.04% of 25(OH)D and 0.4% of 1,25(OH)₂D are free, the remainder being bound to either DBP (85%–88%; high

Figure 7

Relative resistance to vitamin D₃ toxicity demonstrated by DBP⁻/⁻ mice. DBP⁺/⁺ and DBP⁻/⁻ mice were injected with toxic doses of vitamin D or with vehicle alone. (a) Serum calcium levels were determined 7 days after injection, and results are expressed as a percentage of the serum calcium in the vehicle-injected control groups. The differences were significant in both comparisons: DBP⁺/⁺ group (P < 0.001) and DBP⁻/⁻ group (P < 0.01) compared with vehicle (not shown), and the calcium increase in the DBP⁺/⁺ group compared with the DBP⁻/⁻ group (P < 0.05, shown). Seven days after injection, kidney sections were fixed and stained with von Kossa to detect calcium deposits. Representative kidney sections from DBP⁺/⁺ mice injected with vehicle (b) or vitamin D (c), and DBP⁻/⁻ mice injected with vehicle (d) or vitamin D (e) are shown (x10). The arrows (c and e) point to calcium deposits in the renal cortex, and these regions are shown in the insets (x20).
Accelerated activation of calbindin-D_{9k} gene expression by 1,25(OH)_{2}D_{3} in DBP^{-/-} mice. (a) Vitamin D--deficient DBP^{+/+} and DBP^{-/-} mice were injected intravenously with 50 ng 1,25(OH)_{2}D_{3}. Animals were sacrificed at the indicated times after injection, and RNA was isolated from the most proximal centimeter of small intestine. The RNA was analyzed by Northern blots hybridized with 32P-labeled calbindin-D_{9k} and [32P]rpL32 (loading control) probes. A representative autoradiogram is shown. (b) Relative band intensities were quantitated by PhosphorImager and were normalized for RNA loading. The data presented are expressed as a percentage of the maximal intensities were quantitated by PhosphorImager and were normalized for RNA loading. The data presented are expressed as a percentage of the maximal calbindin-D_{9k} mRNA levels in response to 1,25(OH)_{2}D_{3}. The mean ± SEM from four independent experiments is shown.
The very rapid plasma clearance of 25(OH)[3H]D₃ and its greater excretion in the urine in an unmodified form (Fig. 5a) in the absence of DBP suggested that binding of 25(OH)D by DBP slowed its direct filtration and excretion in urine (Fig. 5b). This model was supported by the results of the thin-layer chromatographic fractionation of plasma samples from 0 through 40 minutes after 25(OH)[3H]D₃ injection. These samples demonstrated a stable maintenance of the lipid-soluble tritiated 25(OH)D in DBP⁺/⁻ mice but a rapid fall in DBP⁺/⁻ mice (Fig. 5c). In contrast, polar metabolites increased over 40 minutes in DBP⁺/⁻ mice but fell in DBP⁺/⁻ mice, suggesting that a more rapid and reciprocal conversion of 25(OH)D to biologically inactive polar products was occurring. Thus, the DBP⁺/⁻ mice demonstrated poor conservation of 25(OH)D in serum due to accelerated urinary excretion as well as metabolic inactivation.

In the absence of DBP, [³H]vitamin D₃ entered the liver at an accelerated rate and was cleared from the serum more efficiently than in the wild-type animals (Fig. 6). These results suggested that DBP prolonged the serum half-life of vitamin D and retarded its entry into the liver and its subsequent metabolism. In a single-pass liver-perfusion system, removal of dietary vitamin D was shown to be greatest when it was carried by LDL or chylomicron remnants and thus occurred via receptor-mediated uptake. In contrast, the uptake of vitamin D by hepatocytes was significantly slower when carried on DBP (44). In vivo, some vitamin D is presented to the liver on LDL and chylomicrons and some will be presented on other plasma carriers (DBP, albumin, other lipoproteins), and the overall rate of uptake in the normal animals likely reflects the net internalization of these several carriers (33). It has been previously shown that rapid delivery of vitamin D to the liver leads to the rapid conversion of the excess to inactive forms and water-soluble conjugates by esterification (34). In the experiments reported here, more tritiated serum sterol was in the form of 25(OH)D in the wild type and in the form of polar metabolites in the DBP⁻/⁻ group at one minute after injection. Thus, the rapid entry of vitamin D and its selective shunting to an esterification-excretion pathway would be consistent with the predictions from prior metabolic studies and consistent with this finding. The abnormal levels and conversion of vitamin D to water-soluble side products in the DBP⁻/⁻ mice thus reflected the role of DBP in modulating hepatic uptake of vitamin D and its subsequent 25-hydroxylation and activation. The loss of these functions contributed to the sensitivity of the DBP⁻/⁻ mice to vitamin D deprivation as well as to their relative resistance to vitamin D toxicity.

Vitamin D toxicity is manifested by hypercalcemia, bone resorption, and calcification of soft tissues. In the face of vitamin D toxicity, conversion of vitamin D to the active 1,25(OH)₂D₃ form is normally downregulated, and 25(OH)D or one of its metabolic products has been proposed to be responsible for the direct toxic effect (45, 46). An alternative hypothesis proposes that the toxicity may be secondary to competition from the high levels of 25(OH)D for DBP binding, with resultant elevations of the presumed buffering capacity of DBP in vitamin D metabolism. DBP⁻/⁻ mice were less susceptible to hypercalcemia and the secondary soft-tissue calcification of vitamin D toxicity than the wild-type controls (Fig. 7). Based on the previous results, the relative resistance of DBP⁻/⁻ mice to the full effects of vitamin D toxicity may reflect the net effect of the more rapid direct urinary excretion of 25(OH)D and the more rapid conversion to inactive polar metabolites (Fig. 5). The protective effect of a DBP-null state is also consistent with the previous demonstration that toxic levels of 25(OH)D exceeding the normal levels of DBP’s binding capacity result in an increased fractional liver uptake and increased biliary excretion (48). Furthermore, in the absence of DBP, the competition model would not be active. Thus, the buffering capacity of DBP is deleterious in the face of an abnormally high vitamin D load.

25(OH)D has been demonstrated to enter cells independent of a carrier-mediated transport mechanism in vitro (49). Because of this action and the absence of evidence for functional vitamin D deficiency in DBP⁻/⁻ animals on normal diets, it can be predicted that a fraction of the serum 25(OH)D was taken up by the kidney for 1-hydroxylation in DBP⁺/⁻ mice. Because the serum levels of 25(OH)D in the DBP⁻/⁻ mice were significantly low (Fig. 3), a continuous input of dietary vitamin D would be required to maintain normal 1,25(OH)₂D₃ action in target tissues. This hypothesis is supported by the experimental results, demonstrating a marked sensitivity of the DBP⁻/⁻ mice to dietary deficiency of vitamin D.

Whether sterol-binding proteins facilitate or hinder entry of the corresponding sterol into a target cell remains a critical question. It has been demonstrated that many steroid-binding proteins are associated with cell surface

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Diet (weeks vitamin D-deficient)</th>
<th>Ca⁺⁺ (mg/dl)</th>
<th>P (mg/dl)</th>
<th>Alkaline phosphatase (U/l)</th>
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<tr>
<td>DBP⁺/⁻</td>
<td>0</td>
<td>8.0 ± 0.3</td>
<td>11.4 ± 1.0</td>
<td>74 ± 28</td>
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<tr>
<td></td>
<td>(9)</td>
<td>(9)</td>
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<tr>
<td>DBP⁻/⁻</td>
<td>0</td>
<td>7.9 ± 0.8</td>
<td>10.3 ± 1.5</td>
<td>94 ± 26</td>
</tr>
<tr>
<td></td>
<td>(15)</td>
<td>(16)</td>
<td>(15)</td>
<td></td>
</tr>
<tr>
<td>DBP⁺/⁺</td>
<td>4</td>
<td>8.3 ± 0.6</td>
<td>8.0 ± 1.0⁺</td>
<td>90 ± 31</td>
</tr>
<tr>
<td></td>
<td>(9)</td>
<td>(8)</td>
<td>(8)</td>
<td></td>
</tr>
<tr>
<td>DBP⁻⁻</td>
<td>4</td>
<td>8.3 ± 0.4</td>
<td>8.8 ± 1.2⁺</td>
<td>126 ± 46</td>
</tr>
<tr>
<td></td>
<td>(8)</td>
<td>(8)</td>
<td>(7)</td>
<td></td>
</tr>
<tr>
<td>DBP⁺⁺</td>
<td>6</td>
<td>7.5 ± 1.0</td>
<td>N.D.</td>
<td>119 ± 42⁺⁺</td>
</tr>
<tr>
<td></td>
<td>(14)</td>
<td></td>
<td>(15)</td>
<td></td>
</tr>
<tr>
<td>DBP⁻⁻</td>
<td>6</td>
<td>7.8 ± 0.8</td>
<td>N.D.</td>
<td>131 ± 44⁺⁺</td>
</tr>
<tr>
<td></td>
<td>(13)</td>
<td></td>
<td>(14)</td>
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</table>

For comparisons between vitamin D-deficient and standard diet groups of the same genotype: ⁺P < 0.0001, ⁺⁺P < 0.001, ⁺⁹P < 0.01.
receptors and can be internalized (11, 12, 50). The same has been shown for albumin, a member of the DBP gene family that shares significant amino acid and secondary structural similarities with DBP but that is not a high-affinity steroid/sterol binding protein (4). A variety of cell types have been shown to have DBP on their surfaces through immunohistochemical and immunofluorescent analyses (51–55), suggesting that DBP might mediate entry of vitamin D sterols into the cell. However, in its role as a circulating reservoir for the vitamin D sterols, DBP might alternatively slow the entry of 1,25(OH)2D3 into its target cells by regulating the amount of free sterol available. These possibilities were tested by comparing the kinetics of 1,25(OH)2D3 induction of calbindin-D9K gene expression in vivo. Induction of calbindin-D9K gene expression has served as a sensitive indicator of the biological response of target tissues to 1,25(OH)2D3 action (56, 57). In the mouse, administration of 1,25(OH)2D3 to vitamin D–deficient animals has been shown to induce a fourfold increase in transcription of the calbindin-D9K gene in proximal mouse intestine and a 20-fold induction in the kidney (35). To determine whether the absence of DBP modified the kinetics of calbindin-D9K gene induction by altering the rate at which the hormone entered the target tissue, 1,25(OH)2D3 was injected intravenously and the levels of calbindin-D9K mRNA were monitored in the proximal intestine. The experiment revealed two observations relevant to DBP’s role in gene activation. First, the magnitudes of 1,25(OH)2D3 induction of calbindin-D9K in vitamin D–deficient DBP+/+ and DBP−/− mice were found to be nearly identical (Fig. 8a). Thus, in the absence of DBP, a 1,25(OH)2D3-sensitive gene could still mount a full response to 1,25(OH)2D3 stimulation. The second result was that the kinetic profiles of the response differed (Fig. 8b). The maximum response in DBP+/+ intestine occurred 12 hours after injection, as previously reported (35), whereas the maximal response in DBP−/− proximal intestine occurred at eight hours. This more rapid induction of gene activation could be explained by more rapid entry of the sterol into the target cells of the DBP−/− mice. Thus, the accelerated intracellular response to 1,25(OH)2D3 in the intestinal epithelial cells of the DBP−/− mouse paralleled the more rapid entry of [3H]vitamin D into liver (Fig. 6b).

DBP’s primary role appeared to be the sequestration of vitamin D sterols in the serum, prolonging their serum half-lives and providing a circulating store of 25(OH)D to meet transient periods of vitamin D deficiency. In so doing, DBP helped to prevent the development of severe vitamin D deficiency that leads to bone mineralization defects and the ultimate development of osteomalacia or rickets. As an adjunct to its role in conservation and optimization of vitamin D, DBP appeared to minimize direct urinary losses of the sterols and to slow their entry into metabolic breakdown pathways. One corollary of this phenomenon is that DBP−/− mice, totally lacking DBP, manifested a need for continual intake of vitamin D to avoid the deficiency syndrome and also a reciprocal resistance to vitamin D toxicity in the face of increased sterol load. A testable clinical prediction from these findings is that patients with nephrotic syndrome and decreased serum DBP levels might be more vulnerable to osteomalacia or rickets. It would appear that evolution, in establishing high levels of DBP, has favored the need for conservation and optimal utilization of dietary vitamin D over the more unlikely danger of dietary-induced toxicity.

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