Heat shock protein 47 (HSP47) is an endoplasmic reticulum (ER)-resident collagen-specific chaperone and essential for proper formation of the characteristic collagen triple helix. It preferentially binds to the folded conformation of its clients and accompanies them from the ER to the Golgi compartment, where it releases them and is recycled back to the ER. Unlike other chaperones, the binding and release cycles are not governed by nucleotide exchange and hydrolysis, but presumably the dissociation of the HSP47–procollagen complex is triggered by the lower pH in the Golgi (pH 6.3) compared with the ER (pH 7.4). Histidine residues have been suggested as triggers due to their approximate textbook pKₐ value of 6.1 for their side chains. We present here an extensive theoretical and experimental study of the 14 histidine residues present in canine HSP47, where we have mutated all histidine residues in the collagen binding interface and additionally all of those that were predicted to undergo a significant change in protonation state between pH 7 and 6. These mutants were characterized by bio-layer interferometry for their pH-dependent binding to a collagen model. One mutant (H238N) loses binding, which can be explained by a rearrangement of the Arg²²² and Asp³⁸⁵ residues, which are crucial for specific collagen recognition. Most of the other mutants were remarkably silent, but a double mutant with His²⁷³ and His²⁷⁴ exchanged for asparagines exhibits a much less pronounced pH dependence of collagen binding. This effect is mainly caused by a lower kₐₕ at the low pH values.

Collagen is the most abundant protein in mammals and constitutes a crucial component of the extracellular matrix. There are 28 different types of collagens known today. All of them have in common as a defining feature the so-called collagen triple helix, a structural element that is formed by three individual polypeptide chains (reviewed in Refs. 1–3). The three-dimensional structure of this extended, non-globular domain puts severe restrictions on the linear amino acid sequence of collagens, resulting in repeating triplets of the sequence Xaa-Yaa-Gly with proline and hydroxyproline as the preferred residues in the Xaa and Yaa positions, respectively. Biogenesis of collagen is a complicated process involving extensive post-translational modifications (4). After secretion into the endoplasmic reticulum (ER), a large network of modifying enzymes like proline hydroxylase or sugar transferases as well as molecular chaperones, such as protein-disulfide isomerase or binding immunoglobulin protein, are involved in the formation of a properly modified and folded trimeric procollagen molecule that is secreted via the Golgi compartment to the extracellular medium. HSP47 is an ER-resident procollagen-specific chaperone that binds to multiple sites on procollagen bearing an arginine in the Yaa positions and is essential for proper triple-helix formation (5, 6). HSP47 gene ablation results in impaired secretion, accompanied by overmodification and intracellular procollagen accumulation, as well as defective maturation of the procollagen chains (7). The resulting (pro)collagen from such HSP47-deficient cells exhibits a misfolded triple helix, as indicated by a proteolytic sensitivity of this otherwise very robust structure. As a consequence, the knock-out in mice is lethal at an early embryonic stage due to malformation of the basal membrane (8), and missense mutations in the HSP47/SERPINC1 gene are one cause of recessive osteogenesis imperfecta (9, 10).

Contrary to other chaperones, HSP47 binds preferentially to the folded (i.e. triple-helical) conformation of its client (11), and client binding and release events are not triggered by nucleotide binding and hydrolysis cycles but presumably rather by the pH shift the HSP47–procollagen complex undergoes on its way from the ER (pH ~7.4) to the cis-Golgi compartment (pH ~6.3) (12, 13). After dissociation from collagen in the low pH compartments of the cis-Golgi or ER–Golgi intermediate compartment, HSP47 is recycled back to the ER due to its C-terminal RDEL sequence.

HSP47 is a member of the serpin family (14) but has lost all inhibitory function together with the characteristic serpin S → R transition that results upon proteolytic cleavage or latency transition in the insertion of a mobile element, the so-called reactive center loop, into the middle of a β-sheet. Based
on CD and fluorescence spectra, conformational rearrangements of HSP47 at lower pH values have nevertheless been postulated (15, 16). Crystal structures of HSP47 in complex with collagen model peptides have been published, and the interface between the two proteins is known in atomic detail (17). However, a number of important questions, especially regarding the pH-triggered client release, are still open. Six histidine residues are found within this binding interface, thus opening an attractive explanation of client release at slightly acidic pH values. The involvement of imidazole side chains has also been previously postulated, due to their $pK_a$ values usually falling in the interval between 6 and 7, thus titrating in the pH interval relevant to the HSP47 client-binding and release cycle (15). Previous work has mainly assigned a pair of histidine residues in the so-called breach region of HSP47 as responsible triggers of collagen binding and release (18), together with another histidine located close by. The exact mechanism by which protonation of these histidines results in collagen dissociation from HSP47 is still unclear.

To gain a deeper insight into the function of HSP47 and the molecular nature of the interaction between chaperone and client, we have undertaken a systematic study involving the prediction of $pK_a$ values followed by an extensive mutagenesis approach. We show here that the previous assignment of histidine residues is in error and put forward a detailed model of how low pH triggers collagen release from HSP47.

**Experimental Procedures**

Molecular Dynamics Simulations—The crystal structure of HSP47 in complex with a collagen model peptide (Protein Data Bank entry 3ZHA (17), chains ABFEG) was used as a template for the starting structures. For closing the gap in the protein backbone between Ser-119 and Val-126 (chain A), a modeled loop was inserted using ModLoop (19, 20).

Titrating aspartate, glutamate, and histidine residues were renamed to prepare the pdb files for the constant pH molecular dynamics (CpHMD) simulation developed by Mongan et al. (21). Hydrogen atoms were added using the LEaP module of AMBER 12 (22). Subsequently, the topology and coordinate files of the systems were generated with LEaP. The CpHMD (and also all other MD) simulations were performed using Sander of AMBER 12 and the ff99SB force field (23) for proteins. At first, a minimization was carried out in three subsequent steps to optimize the geometry of the starting structures. In the first step of the minimization, the hydrogen atoms of the protein were minimized, whereas everything else was restrained with a constant force of 10 kcal/(molÅ$^2$) to the initial positions. In the second step, the modeled loop was minimized, whereas the remaining protein was restrained with 10 kcal/(molÅ$^2$). In the last step, no restraints were used, so that the whole system was minimized. All three minimization parts started with 2,500 steps using the steepest descent algorithm, followed by 2,500 steps of a conjugate gradient minimization.

The molecules were solvated with the implicit solvent developed by Onufriev et al. (control variable igb = 2) (24). The salt concentration was set to 0.1m (based on Debye–Hückel). The cut-off distance for non-bonded interactions was set to 30.0 Å, and bonds involving hydrogen were constrained with SHAKE. The time step was 2 fs, and a Monte Carlo step was performed after every step. Thereafter, the systems were equilibrated in two successive steps. In the first step (0.1 ns long), the temperature was raised from 10 to 310 K, and the protein was restrained with a constant force of 5 kcal/(molÅ$^2$). In the second step (0.4 ns long), only the Ca atoms of the protein were restrained with a constant force of 5 kcal/(molÅ$^2$). Then the simulations ran for 25 ns (pH 4.0) or 100 ns (pH 5.0, 6.0, 7.0, and 8.0) without any restraints at physiological temperature (310 K) with a Langevin thermostat. For the analysis of the MD simulations, the fraction of doubly protonated histidines was measured over simulation.

The H238N mutant and, for comparison, also the wild-type HSP47 were simulated without the collagen molecule for 50 ns with the described CpHMD method at pH 8.0. For the wild-type HSP47 and the mutants (H273N, H274N, and H273N/H274N), a modified version of the CpHMD, namely a pH titrating MD (pHtMD) (25) simulation, was conducted. During these 201-ns-long simulations, the solution pH was decreased from 8.0 to 4.0 with a rate of $-0.02$ pH units/ns.

Steered molecular dynamics (SMD) simulations were performed for wild-type HSP47 and the H273N/H274N mutant, both complexed with the collagen model peptide, to investigate the work necessary for dissociation of the HSP47-collagen complex. Ten different structures from the pHtMD simulation at pH 6.0 were used as starting structures for the 10 independent SMD simulations. While the HSP47 molecule was restrained with a constant force of 5,000 kcal/(molÅ$^2$), the collagen was pulled away with a constant velocity of 0.5 Å/ps, thereby increasing the distance between the centers of mass of the collagen triple helix and the HSP47 molecule from 32 to 82 Å. Finally, the computed work, which is the integrated force over distances, was averaged over the 10 simulations.

Protein Constructs and Expression—A canine-derived synthetic HSP47 gene coding for residues $^{36}$LSP...$^{36}$DEL with a modified version of the CpHMD, namely a pH titrating MD (pHtMD) (25) simulation, was conducted. During these 201-ns-long simulations, the solution pH was decreased from 8.0 to 4.0 with a rate of $-0.02$ pH units/ns.

Steered molecular dynamics (SMD) simulations were performed for wild-type HSP47 and the H273N/H274N mutant, both complexed with the collagen model peptide, to investigate the work necessary for dissociation of the HSP47-collagen complex. Ten different structures from the pHtMD simulation at pH 6.0 were used as starting structures for the 10 independent SMD simulations. While the HSP47 molecule was restrained with a constant force of 5,000 kcal/(molÅ$^2$), the collagen was pulled away with a constant velocity of 0.5 Å/ps, thereby increasing the distance between the centers of mass of the collagen triple helix and the HSP47 molecule from 32 to 82 Å. Finally, the computed work, which is the integrated force over distances, was averaged over the 10 simulations.

Protein Constructs and Expression—A canine-derived synthetic HSP47 gene coding for residues $^{36}$LSP...$^{36}$DEL was purchased as a codon-optimized cDNA for Escherichia coli from DNA 2.0 (Menlo Park, CA) with an additional C-terminal StrepII tag and cloned into the manufacturer’s expression plasmid, pExpress 411. The construct was transformed in E. coli BL21 (DE3) cells grown to $A_{\text{om}}$ of 0.6–0.7, and protein expression was induced with 0.5 mm isopropyl-$\beta$-d-thiogalactoside. After a 4-h incubation at 37 °C/180 rpm, cells were harvested, and pellets were stored at −20 °C. Cells were resuspended in lysis buffer (50 mm Tris HCl (pH 7.5), 150 mm NaCl, 100 μm PMSF, 4 mm DTT) and lysed by sonication. Cleared lysates were loaded onto 1 ml of Streptactin Superflow (IBA) and eluted with 2.5 mm d-Desthiobiotin in lysis buffer excluding PMSF. The eluted proteins were further subjected to size exclusion chromatography (Superdex 200, GE Healthcare) in 50 mm Hepes (pH 7.5), 150 mm NaCl, 2 mm DTT, and the protein was frozen in liquid nitrogen and stored at −80 °C.

Mutations were introduced according to an optimized site-directed mutagenesis protocol (26) using overlapping primers. The mutants were expressed and purified following the protocol used for wild-type HSP47.

The plasmid coding for the GPR-foldon peptide (27) was a generous gift from Dr. M. Koch (University of Cologne). It encodes an N-terminal Twin-StrepII tag, a thrombin site, a collagen sequence ((GPP)$_5$GPR(GPP)$_6$), and a C-terminal phage
T4 foldon domain for protein trimerization. The protein was expressed in BL21 (DE3), induced at $A_{600}$ of 1.0 with 100 $\mu$M isopropyl-$\beta$-D-thio-galactoside, and incubated overnight at 20 °C/180 rpm. The foldon protein was purified analogously to HSP47 constructs but without DTT in the buffers.

**Phylogenetic Analysis**—Protein sequences of all currently annotated SerpinH1 orthologs in ENSEMBL (ensemble.org/GRCh38.p3) were retrieved and aligned using the ClustalO plugin of CLC Bio. Sequences containing too many gaps to be useful were deleted. The species were grouped together based on their phylogenetic class, and these classes were ordered by their phylogenetic relationship. Protein sequences of human SerpinF1 (PEDF), SerpinA1 (antitrypsin), and SerpinC1 (antithrombin) were included as an outgroup. The alignment was colored using BOXSHADE.

**Kinetic Measurements Using Biolayer Interferometry**—Biolayer interferometry was measured on a BLItz system (Pall ForteBio LLC, Menlo Park, CA). For coupling on streptavidin biosensors, foldon peptides were biotinylated with NHS-Biotin following the manufacturer’s protocols (Pierce) and loaded at a concentration of 10 $\mu$M to a response of $\sim$5.5 nm. Free biotin binding sites were blocked with 10 $\mu$M biocytin, and nonspecific interactions were blocked with two load/regeneration cycles with HSP47. The sensors were regenerated with McIlvaine buffer (28) at pH 6.0 for up to 40 s. Successful regeneration of the sensors was verified by comparison of HSP47 binding at pH 7.5 before and after regeneration. Kinetics were measured with a constant protein concentration of 5 $\mu$M HSP47 protein. The association time was fixed at 60 s, whereas the dissociation time was chosen differently at different pH to allow for sufficient dissociation (pH 7.5, 140 s; pH 7.0, 120 s; pH 6.5, 100 s; pH 6.0, 80 s). For each condition and sample, at least three sensograms were recorded, and all experiments were repeated at least twice on different days with different protein preparations. Kinetic parameters were extracted separately using the program Origin version 8.5 and a 1:1 Langmuir binding model. Sensograms are shown as representative results, and kinetic parameters ($k_{on}$, $k_{off}$, and $K_D$) are presented with error bars indicating the S.D. in a single experiment.

**ELISA Style Binding Assays**—For ELISA style binding assays, rat tail collagen I (BD Biosciences) was diluted in TBS and coated at 10 $\mu$g/ml (500 ng/well) overnight at room temperature onto 96-well plates (MaxiSorp, Nunc). After washing with TBS-T (TBS containing 0.05% (v/v) Tween 20), plates were blocked for 1 h at room temperature with 5% (w/v) milk powder in TBS. Strep-tagged HSP47 proteins were added at a concentration of 3 $\mu$M in TBS-T containing 1% milk powder and incubated for 1 h. After washing away unbound protein with TBS-T, wells were incubated with McIlvaine buffers at the indicated pH levels for 5 min, shortly washed at the indicated pH, and then washed three times with TBS-T. Bound proteins were detected by Streptactin-HRP (IBA, Goettingen, Germany) and visualized with tetramethylbenzidine as substrate. The reaction was monitored at 370 nm. Data were analyzed using Origin version 8.5 and fitted using a four-parameter logistic model (29).

**CD Spectra**—Circular dichroism spectra were recorded using a Jasco J-715 instrument at 20 °C. All constructs were dialyzed against PBS, pH 7.0, and the indicated pH was adjusted shortly before the measurement by diluting the protein to a final concentration of 0.1 mg/ml in at least 10 volumes of target buffer. The samples were measured in a 1-mm cuvette with bandwidth at 2 nm, a response of 2 s, sensitivity set to standard (100 millidegrees), data pitch 0.2 nm, and a scan speed of 50 nm/min. Each curve was measured with 10 accumulations and plotted in Origin version 8.5. Data were analyzed between 200 and 240 nm with the CDPro package (30) using the CONTIN/LL algorithm (31) and the reference set 7 (30).

**Thermal Shift Assays**—Melting temperatures were measured using a total volume of 50 $\mu$l with 0.1 mg/ml protein concentration and a final dilution of 1× Sypro Orange (Molecular Probes). Samples were heated in a CFX96 real-time PCR system (Bio-Rad) at an approximate rate of 0.5 K/min, and the fluorescence intensity was measured every 0.5 K using the channel 6 filter set (excitation, 450–490 nm; emission, 560–580 nm). Curves were analyzed using the supplied software CFX Manager version 3.1, and melting temperatures were deduced from the minima of the first derivative.

**Results**

**Theoretical Analysis of $pK_a$ Values**—The crystal structure of HSP47 bound to trimeric collagen model peptides (17) (Fig. 1) enabled a structure-based analysis of the pH-dependent client release of HSP47 in the Golgi compartment. During the secretion process, the pH changes from 7.4 in the ER to 6.3 in the Golgi. It was already proposed earlier that histidine residues, whose imidazole side chains have a $pK_a$ of $\sim$6.1, might be responsible for the dissociation of the complex at low pH (15, 18). Canine HSP47 possesses 14 histidine residues (Fig. 1), with most of them being conserved in the HSP47 family. His$^{108}$ and
His^{353} appear to be general serpin features because they occur in all other serpin clades as well, such as antitrypsin (SerpinA1), antithrombin (SerpinC1), or PEDF (SerpinF1). Some histidines show a strict conservation in HSP47 (e.g. His^{238} or His^{315}), whereas others are sometimes replaced (e.g. His^{216}, His^{273}, or His^{362}) (Fig. 2). Of the 14 histidines in HSP47, six are directly

**FIGURE 2. Conservation of titratable histidines in HSP47 orthologs.** Protein sequences from all annotated SerpinH1 orthologs currently in ENSEMBL (ensemble.org/GRCh38.p3) were deduced and aligned using ClustalO. Sequence fragments containing histidines either in the HSP47-collagen interface or predicted to be of importance for pH dependence are shown, separated by white bars. The species were grouped together based on their phylogenetic class, and these classes were ordered by their phylogenetic relationship. Protein sequences of human SerpinF1 (PEDF), SerpinA1 (antitrypsin), and SerpinC1 (antithrombin) were included to differentiate between conservation within HSP47 orthologs and other members of the serpin protein family.
involved in the collagen binding interface, namely His215, His216, His238, His273, His274, and His386 in the numbering of the protein from *Canis lupus familiaris*, UniProt ID C7C419 (Fig. 1). However, histidine residues outside of the interface could introduce changes in the overall structure upon protonation as well and might thereby also lead to a dissociation of the complex. Large conformational changes are very common for members of the serpin family, and pH-dependent changes in CD and fluorescence spectra have been reported (32).

The pK\textsubscript{a} values of histidine side chains are known to depend on the chemical environment generated by the protein structure and can vary from about 4 to 9 with values between 5.0 and 8.0 being regularly observed (33). To narrow down possible key residues, we tried to calculate the pK\textsubscript{a} values of all histidine residues in HSP47 by CpHMD simulations.

These simulations were performed for the HSP47 molecule in complex with the collagen triple helix at pH 8.0, 7.0, 6.0, 5.0, and 4.0, based on the assumption that these histidine residues should change their charge in a pH range between 6.0 and 7.0. Therefore, the doubly protonated (i.e. positively charged) fraction of each of the 14 histidine residues was calculated with an analysis tool of Amber 12 (Fig. 3 and Table 1).

Only 5 of the 14 histidine residues show a significant increase (\textgreater;25\%) in the doubly protonated (i.e. positively charged) state upon decreasing pH from 7.0 to 6.0 (Table 1 (boldface type) and Fig. 3). Only one of these five histidine residues (His273) is
Located close to the interface (Fig. 1, bottom right, red residue). Interestingly, His$_{386}$, a direct neighbor of the important Asp$_{385}$, is already doubly protonated at pH 8.0 and keeps this positive charge over the whole pH range, indicating a strong increase of its pK$_a$ value. This is presumably caused by the neighboring negative charge of Asp$_{385}$, which stabilizes this doubly protonated form of His$_{386}$.

Site-directed Mutagenesis and Preparation and Characterization of HSP47 Mutants—Assuming that the protonation of one or more histidine residues could trigger client release, we generated a set of mutants by changing histidines to asparagines with the rationale that the side chain of asparagine is structurally most similar to the imidazole moiety but cannot be protonated. Based on the simulations and our crystal structure, we replaced every histidine located within the binding interface as well as all of those predicted to change their protonation state by more than 25% between pH 6.0 and 7.0. With the exception of the mutants H386N and H315N, all proteins expressed as well as the wild type and generally exhibited a very similar thermostability, as judged by thermal shift assays (Table 2). The mutant H238N expressed well but had a significantly lower thermal stability, as will be discussed below.

Wild-type HSP47 Releases Its Client Due to a Larger $k_{off}$ at Low pH—We first measured the interaction after washing with buffers of different pH values in an ELISA style binding assay utilizing coated rat tail collagen I. As expected, a decrease in the amount of bound HSP47 can be observed below a pH of 7.0, with a midpoint at a pH of ~5.9 (Fig. 4). To examine this more closely, we investigated the kinetic parameters of the HSP47 interaction with collagen using biolayer interferometry (Fig. 5). When incubated at constant HSP47 concentrations but under different pH values, there was a reduced binding at lower pH to the faldon collagen model peptide (Fig. 5, first panel, WT). Interestingly, the $k_{off}$ rate constants were more strongly affected than the $k_{on}$ values. For the wild type, the $k_{off}$

### TABLE 1
Protonation states of histidines in HSP47

| Residue number | Interface | Fraction of doubly protonated histidine residues |
|---------------|-----------|-----------------------------------------------|
|               | pH 4.0 | pH 5.0 | pH 6.0 | pH 7.0 | pH 8.0 |
| 108           | 94.9   | 92.9   | 14.1   | 5.0    | 0.1    |
| 153           | 95.8   | 90.3   | 67.1   | 6.2    | 1.3    |
| 158           | 96.1   | 62.4   | 36.9   | 5.8    | 1.0    |
| 209           | 79.6   | 59.4   | 11.1   | 10.4   | 0.4    |
| 215           | Yes    | 5.6    | 3.7    | 0.1    | 0.2    |
| 216           | Yes    | 94.8   | 75.7   | 15.8   | 0.5    |
| 238           | Yes    | 42.3   | 4.9    | 1.7    | 1.4    |
| 262           | 99.0   | 90.0   | 82.6   | 18.9   | 3.1    |
| 273           | Yes    | 91.6   | 50.3   | 32.7   | 3.3    |
| 274           | Yes    | 20.9   | 5.4    | 0.5    | 0.1    |
| 315           | 50.4   | 83.7   | 59.3   | 11.1   |
| 320           | 87.8   | 9.2    | 11.7   | 13.4   |
| 353           | 80.5   | 67.5   | 4.5    | 23.9   |
| 386           | Yes    | 100.0  | 100.0  | 97.9   | 98.0   |

**FIGURE 4.** The pH dependence of client binding agrees with the titration curves of histidine residues. There is a clear pH dependence in an ELISA style HSP47-collagen binding assay employing rat tail collagen I as coat and recombinant canine HSP47 as analyte, indicating an inflection point around pH 5.88 ± 0.03 for the WT protein. The double mutant H273N/H274N shows a decreased inflection point at pH 5.66 ± 0.02. Data points were measured in triplicates, and the results were confirmed using at least two independent biological samples. Error bars, S.D.

### TABLE 2
Melting temperatures and dissociation rate constants ($k_{off}$) of HSP47 variants

| Protein | $T_m$ °C | $k_{off}$ $s^{-1}$ | pH 6.0 | pH 6.5 | pH 7.0 | pH 7.5 |
|---------|----------|-------------------|--------|--------|--------|--------|
| Wild type | 57 | 0.192 ± 0.002 | 0.075 ± 0.001 | 0.045 ± 0.001 | 0.029 ± 0.001 |
| H153N | 56 | 0.179 ± 0.003 | 0.074 ± 0.006 | 0.045 ± 0.001 | 0.031 ± 0.003 |
| H158N | 56 | 0.187 ± 0.002 | 0.070 ± 0.002 | 0.045 ± 0.001 | 0.031 ± 0.003 |
| H215N | 58 | 0.313 ± 0.023 | 0.159 ± 0.015 | 0.126 ± 0.005 | 0.097 ± 0.005 |
| H216N | 57 | 0.184 ± 0.013 | 0.088 ± 0.004 | 0.055 ± 0.003 | 0.039 ± 0.001 |
| H215N/H216N | 58 | 0.273 ± 0.005 | 0.168 ± 0.019 | 0.124 ± 0.003 | 0.094 ± 0.002 |
| H238N | 45 | ND | ND | ND | ND |
| H262N | 57 | 0.236 ± 0.018 | 0.096 ± 0.002 | 0.047 ± 0.004 | 0.030 ± 0.001 |
| H273N | 55 | 0.148 ± 0.009 | 0.063 ± 0.003 | 0.038 ± 0.003 | 0.028 ± 0.001 |
| H274N | 56 | 0.170 ± 0.010 | 0.075 ± 0.004 | 0.038 ± 0.001 | 0.029 ± 0.001 |
| H273N/H274N | 56 | 0.071 ± 0.002 | 0.038 ± 0.002 | 0.026 ± 0.001 | 0.024 ± 0.001 |
| H273A | 57 | 0.111 ± 0.002 | 0.048 ± 0.004 | 0.030 ± 0.002 | 0.023 ± 0.002 |
| H274A | 57 | 0.169 ± 0.005 | 0.071 ± 0.008 | 0.036 ± 0.002 | 0.029 ± 0.001 |
| H273A/H274A | 57 | 0.080 ± 0.002 | 0.043 ± 0.002 | 0.031 ± 0.001 | 0.028 ± 0.002 |
| H273D | 57 | 0.110 ± 0.003 | 0.051 ± 0.002 | 0.028 ± 0.001 | 0.020 ± 0.001 |
| H273K | 58 | 0.134 ± 0.006 | 0.057 ± 0.005 | 0.034 ± 0.001 | 0.023 ± 0.001 |
| H274D | 58 | 0.100 ± 0.002 | 0.048 ± 0.004 | 0.032 ± 0.002 | 0.024 ± 0.001 |
| H274K | 57 | 0.285 ± 0.011 | 0.167 ± 0.006 | 0.124 ± 0.012 | 0.103 ± 0.010 |
| H273F | 50 | 0.139 ± 0.003 | 0.050 ± 0.002 | 0.029 ± 0.000 | 0.022 ± 0.000 |
| H273L | 57 | 0.112 ± 0.002 | 0.039 ± 0.001 | 0.024 ± 0.001 | 0.017 ± 0.000 |
| H273Y | 57 | 0.215 ± 0.001 | 0.076 ± 0.002 | 0.043 ± 0.003 | 0.029 ± 0.001 |

* a Mean values for all experiments.
pH Dependence Mechanism of the HSP47-Collagen Interaction

WT

H153N

H158N

H215N

H216N

H215N:H216NN

H262N

H273N

H274N

H273N:H274N

H273A

H274A

HH273:274AA
increased from 0.028 s\(^{-1}\) at pH 7.5 to 0.192 s\(^{-1}\) at pH 6.0 (Table 2), similarly to the \(K_D\) change from about 0.74 to 6.23 \(\mu M\) (Fig. 6 (WT) and Table 3). This change is mainly driven by the increase in \(k_{\text{off}}\) because the \(k_{\text{on}}\) stays relatively constant at about 40,000 \((M\cdot s)^{-1}\) and 33,000 \((M\cdot s)^{-1}\) (Fig. 6). Interestingly, the kinetic parameters of all of the mutants not located within the binding interface (H153N, H158N, and H262N) were similar to those of the wild-type protein (Figs. 3 and 6 and Tables 2 and 3), thus excluding a long-range conformational rearrangement upon protonation.

His\(^{238}\) Is Essential for Collagen Binding by Organizing an Extended Hydrogen Bond Network but Does Not Govern the pH Switch—

His\(^{238}\) is located in the center of the interaction surface between collagen and HSP47. The single mutant H238N has no detectable binding to collagen anymore (Fig. 7A). The side chain of His\(^{238}\) forms few and rather weak contacts with the collagen...
In the wild-type situation, His\textsuperscript{238} is able to form hydrogen bonds with the side chains of Asp\textsuperscript{220} and Ser\textsuperscript{305} (Fig. 7B, dashed lines). Asp\textsuperscript{220} forms an additional stable salt bridge to the Ne and N\texteta\textsuperscript{2} atoms of Arg\textsuperscript{222} (Fig. 7B, blue dashed lines), thus, together with an additional hydrogen bond from Tyr\textsuperscript{383}, stabilizing this side chain in a conformation capable of binding to the collagen backbone (Fig. 7B, dashed lines). Calculation of the protonation states (Fig. 3) indicates that His\textsuperscript{238} starts to titrate only at pH < 5; thus, it should not govern the pH switch. Its shifted pK\textsubscript{a} value can be attributed to the adjacent positive charge of the Arg\textsuperscript{222} side chain, which hampers protonation at the Ne nitrogen.

To understand the effect of the H238N mutation, we modeled it in collagen-unbound HSP47 and simulated the conformational changes by constant pH molecular dynamics simulations. For comparison, additional simulations of collagen-bound and unliganded wild-type HSP47 were performed. Monitoring the Asp\textsuperscript{220}-Arg\textsuperscript{222} side-chain interaction over the simulation time reveals that this interaction is both present in the collagen-bound and in the unbound wild-type HSP47 (Fig. 7D, gray and black). This implies that the respective interaction plays an important role in prepositioning Arg\textsuperscript{222} for a proper interaction with collagen. In contrast, this interaction is significantly less stable in the unliganded H238N mutant (Fig. 7D,

The table shows the dissociation constants (K\textsubscript{D}) of HSP47 variants. The values are means ± S.D. ND, not determined.

| Protein            | pH 6.0 | pH 6.5 | pH 7.0 | pH 7.5 |
|--------------------|--------|--------|--------|--------|
| Wild type          | 6.23 ± 0.08 | 2.25 ± 0.24 | 1.14 ± 0.04 | 0.74 ± 0.02 |
| H153N              | 6.06 ± 0.21 | 1.93 ± 0.46 | 1.20 ± 0.04 | 0.76 ± 0.02 |
| H158N              | 3.85 ± 0.52 | 1.38 ± 0.13 | 1.00 ± 0.08 | 0.55 ± 0.05 |
| H215N              | 8.93 ± 1.20 | 3.70 ± 0.96 | 2.21 ± 0.27 | 1.60 ± 0.19 |
| H216N              | 7.65 ± 1.93 | 2.76 ± 0.04 | 1.55 ± 0.14 | 1.18 ± 0.08 |
| H215N/H216N        | 4.58 ± 0.92 | 4.70 ± 1.96 | 2.15 ± 0.12 | 1.63 ± 0.12 |
| H238N              | ND      | ND      | ND      | ND      |
| H262N              | 11.39 ± 5.18 | 2.55 ± 0.17 | 1.13 ± 0.20 | 0.67 ± 0.06 |
| H273N              | 7.27 ± 2.01 | 2.34 ± 0.29 | 1.27 ± 0.13 | 0.97 ± 0.04 |
| H274N              | 5.86 ± 0.75 | 2.12 ± 0.26 | 0.92 ± 0.05 | 0.65 ± 0.03 |
| H273N/H274N        | 2.60 ± 0.08 | 1.26 ± 0.09 | 0.86 ± 0.02 | 0.85 ± 0.09 |
| H273A              | 6.21 ± 1.20 | 2.58 ± 0.16 | 1.48 ± 0.21 | 0.82 ± 0.09 |
| H274A              | 6.85 ± 1.93 | 2.37 ± 0.19 | 1.12 ± 0.05 | 1.23 ± 0.12 |
| H273A/H274A        | 3.17 ± 0.92 | 1.69 ± 0.23 | 1.00 ± 0.06 | 1.07 ± 0.07 |
| H273D              | 4.04 ± 0.58 | 1.74 ± 0.07 | 0.89 ± 0.02 | 0.64 ± 0.04 |
| H273K              | 5.08 ± 0.67 | 2.11 ± 0.56 | 0.91 ± 0.06 | 0.69 ± 0.02 |
| H274D              | 4.02 ± 0.44 | 1.40 ± 0.07 | 1.11 ± 0.18 | 0.73 ± 0.04 |
| H274K              | 16.29 ± 7.56 | 5.17 ± 0.81 | 3.40 ± 0.78 | 2.98 ± 0.28 |
| H273F              | 5.39 ± 0.20 | 2.20 ± 0.32 | 1.01 ± 0.20 | 0.61 ± 0.04 |
| H273L              | 3.84 ± 0.26 | 1.28 ± 0.06 | 0.72 ± 0.06 | 0.47 ± 0.06 |
| H273Y              | 10.37 ± 1.90 | 3.16 ± 0.20 | 1.53 ± 0.24 | 1.02 ± 0.02 |
**PH Dependence Mechanism of the HSP47-Collagen Interaction**

**The Double Mutant H273N/H274N Shows an Altered PH Dependence of Collagen Binding**—Of all of the mutants that could be studied, only H238N, which was discussed above, and the tandem histidine pair variants H215N/H216N and H273N/H274N showed a significant change in their kinetic behavior. These two tandem histidine pairs are located at opposite ends of the binding interface (Fig. 1).

The H215N/H216N double mutant has been reported to show a loss of the pH switch in binding to collagen when the pH-dependent elution profiles of wild type and mutant in collagen affinity chromatography are compared (18). Therefore, we investigated these residues in more detail by biolayer interferometry. Whereas the H216N variant behaved similar to the wild type in our experiments, the H215N mutant showed an increased k_off over the whole pH range (0.097–0.313 s⁻¹) (Figs. 5 and 6A). Consequently, the double mutant H215N/H216N also showed an overall increase in k_off.

In contrast, the single mutations H273N and H274N individually did not show a significant change compared with the wild-type, whereas the double mutant H273N/H274N showed a markedly reduced pH dependence of the k_off rate constant (Figs. 5 and 6A). In other words, the k_off of the double mutant does not increase as much at lower pH values as the k_off of the wild-type protein or the other histidine mutants that we examined in this study. This is also reflected in the K_ph values of this mutant at the different pH values, which only increase from 0.85 μM at pH 7.5 to 2.60 μM at pH 6.0. This is a significant reduction compared with the wild-type situation (0.74–6.23 μM) (Fig. 6C and Table 3). The tighter binding of the H273N/H274N mutant at pH 6.0 could also be observed in steered molecular dynamics simulations of the dissociation process. Here, more energy was needed to displace the collagen helix from the H273N/H274N double mutant than from the wild-type protein (Fig. 8).

To test whether this effect is specific for the histidine to asparagine exchange, we also replaced these histidine residues with alanines. Biolayer interferometry analysis of the double mutant H273A/H274A showed a behavior similar to that of the asparagine double mutant (Figs. 5 and 6A). However, a small decrease of the k_off at lower pH could already be observed with the single mutant H273A, indicating that this type of exchange might be slightly more important for the pH dependence of the k_off.

The H273N/H274N Double Mutant Is Structurally Similar to the Wild-Type Protein—Based on CD and fluorescence spectra, it was reported earlier that HSP47 undergoes structural changes at low pH (15, 16). To investigate whether the H273N/H274N double mutant has any influence on the overall structure of HSP47, we measured far-UV CD spectra of the mutant and the wild-type protein at pH 7.5 and pH 6.0 (Fig. 9). For the
We noted that the titration of His\textsuperscript{274} in the H273N mutant still starts at lower values than one would expect for a residue that participates in pH-dependent collagen release. This finding can most likely be attributed to the limited accuracy of the modeled mutant structure used for pK\textsubscript{a} value calculations, which are highly sensitive to the local geometry. Nevertheless, the differences between the wild type and H273N mutant are so large that they at least allow the qualitative statement that the identity of the residue at position 273 significantly affects His\textsuperscript{274} titration behavior.

Introducing Charged Residues at Positions 273 and 274—To further test whether charges at the positions of the tandem histidine pair 273/274 impair collagen binding, we mutated these residues to lysine and aspartic acid. Mutation of either histidine to aspartic acid resulted in a slower dissociation at lower pH of the HSP47-client complex (Fig. 11), thus emphasizing again the importance of a positive charge at these positions for client release. As expected, the H274K mutant showed a larger k\textsubscript{off} over the whole pH range observed (i.e. also at neutral pH (pH 7.0–7.5), where the wild type has presumably not acquired yet a significant positive charge at this position). However, this mutant still shows a very pronounced further increase of k\textsubscript{off} at a lower pH. Unexpectedly, the H273K mutant exhibited a decreased dependence of its k\textsubscript{off} on the pH value (Fig 11B, blue curve). A possible explanation is the influence of residue 273 on the pK\textsubscript{a} of residue 274; a positive charge could potentially impede protonation of His\textsuperscript{274}, whereas the flexibility of the lysine side chain and its position further away from the interface as compared with residue 273 could avoid interference with client binding.

Phylogenetic Analysis Confirms the Importance of His\textsuperscript{215}, His\textsuperscript{238}, and His\textsuperscript{274} and Implies Certain Restrictions on His\textsuperscript{273} Replacements—To investigate the conservation of the identified histidines on HSP47, we analyzed all currently listed orthologs of human SerpinH1 in the Ensembl database (ensemble.org/Ch38.p3). As described for many other genes, SerpinH1 exists in two copies in most fish (class Actinopterygii) due to a genome-wide duplication in their common ancestor (34, 35). For medaka and fugu, however, no second gene is currently annotated. Genes originating from whole genome duplication are called ohnologs and may differ in function. Interestingly, many residues in HSP47 are strictly conserved in both fish ohnologs (e.g. MMHRT cluster around His\textsuperscript{238}; Fig. 2), indicating that many of these genes still fulfill a functional role. Most of the histidine residues are relatively well conserved. However, some ohnologs have accumulated exchanges of otherwise well conserved histidines (compare cod genes; Fig. 2, blue boxes).

We have identified four histidine residues that have an effect on binding of HSP47 to collagen. His\textsuperscript{238} (Fig. 2, blue residues), which is essential for binding to collagen, is strictly conserved throughout all species. The tandem pair His\textsuperscript{215}/His\textsuperscript{216} is only conserved in tetrapods (mammals, amphibia, reptiles, and birds) with the exception of the platypus (Fig. 2, green residues), where the correct protein sequence has not yet been confirmed, though. In fish (Actinopterygii and coelacanth) His\textsuperscript{215} is mostly conserved with the exception of one ohnolog in the stickleback. His\textsuperscript{216}, however, is replaced by various residues in a number of genes and also in two ohnologs of one species simultaneously (e.g. platyfish, cod, and tilapia), indicating that this residue is not important for proper HSP47 function at least in these species.
The tandem His\textsuperscript{273}/His\textsuperscript{274} shows a similar behavior. In mammals, both residues are completely conserved (Fig. 2, red residues). In reptiles and birds (sauropsida), however, these residues are exchanged to an asparagine/histidine motif (NH). Because the amphibia still retain an HH motif, the NH of reptiles and birds is very likely to be evolutionarily derived, whereas the two histidines (HH) appear to be more ancestral. In ray-finned fish (Actinopterygii), YH is the most common sequence motif (14 of 18 species) and therefore is likely to be ancestral to ray-finned fish. It is difficult to judge which of the two motifs, HH or YH, is ancestral to Actinopterygii and Sarcopterygii. There are appearances of both motifs in both clades (e.g. YH in coelacanth and HH in platyfish). The more ancestral lamprey carries an LH motif at this position. LH and HH are interchangeable by a single base substitution on the DNA level (changing Leu for His), whereas the mutation of LH to YH requires two bases to be exchanged. This slightly favors an HH motif either as ancestral to all vertebrates (and the lamprey sequence would be a derivative) or as a derived motif from an ancestral LH motif. In summary, substitution of His\textsuperscript{273} by an asparagine or a tyrosine seems not to interfere with HSP47 function. In contrast, His\textsuperscript{274} is evolutionarily more strictly conserved.

A Tyrosine Residue Is Equally Tolerated at the Position of His\textsuperscript{273}—To confirm our theory based on the phylogenetic analysis, we simulated the situation in fish by changing in the canine
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HSP47, the position His\(^{273}\) to tyrosine (most fish), leucine (lamprey and one ohnolog in cod), and phenylalanine (one ohnolog in spotted gar) (Fig. 12). Interestingly, H273Y behaved exactly as the wild-type protein, indicating that the tyrosine at the position 273 is equally able to shift the p\(_K_a\) value of His\(^{274}\) to higher values. In contrast, H273F showed only a small increase in the dissociation rate constants from 0.022 s\(^{-1}\) at pH 7.5 to 0.139 s\(^{-1}\) at pH 6.0 (Fig. 12, blue line) and Table 2). This is similar to the effect of the H273A variant. The H273L showed a significantly reduced pH dependence similar to the H273N/H274N double mutant (Fig. 12, gray line), although at pH 6, the \(k_{\text{off}}\) is slightly larger.

**Discussion**

Our recent crystal structure allowed for the first time the analysis of all histidine residues in HSP47 in their spatial context. For a histidine residue to play a major role in the pH-dependent release, it should change its protonation state from single to doubly protonated between pH 7.0 and 6.0. A total of 14 histidines are present in *C. lupus familiaris* HSP47, of which only 5 are predicted to undergo significant changes in protonation (His\(^{153}\), His\(^{158}\), His\(^{262}\), His\(^{273}\), and His\(^{315}\)). Other, especially buried, histidines are not predicted to get significantly protonated even at pH 6.0.

We reasoned that an exchange of histidines to a non-protonatable analogue should prevent the pH-dependent release. From the 20 canonical amino acids, asparagine is considered to be structurally the closest; it carries a nitrogen in the form of an NH at a similar position as the N\&1 atom of histidine but lacks a further proton acceptor similar to the Ne2 atom. From our analysis of the wild-type interaction, we know that the pH dependence is mostly conveyed by a increase in \(k_{\text{off}}\) between pH 7 and pH 6. We therefore exchanged all histidines that are predicted to undergo a large change in protonation state as well as all the histidines in the interface to asparagine residues and measured their kinetic parameters.

Two of these HSP47 mutants (H315N and H386N) did not express in a soluble form. This can be explained by the fact that both residues are buried in the structure and are probably important for the integrity of the folded conformation.

H238N did not show any binding to collagen. This residue is located in a catalytic triad-like arrangement with Ser\(^{305}\) and Asp\(^{220}\). Analysis of this mutant in molecular simulations showed that His\(^{238}\) is critical for the stabilization of a hydrogen network locking Arg\(^{222}\) in a binding-capable conformation, thus preventing an intramolecular salt bridge between Arg\(^{222}\) and Asp\(^{385}\), two residues of pivotal importance for the HSP47 collagen interface.

In an earlier study employing recombinant mouse HSP47, His\(^{191}\), His\(^{197}\), and His\(^{198}\) (numbering according to the mature mouse protein), which correspond to His\(^{209}\), His\(^{215}\), and His\(^{216}\) (numbering according to the full-length canine HSP47 as used here) were reported to be important for the pH-dependent client release. It was observed that a double mutant corresponding to H215N/H216N and a single mutant corresponding to H209A were released early in a pH elution from a collagen affinity column. H215N/H216N showed a particularly broad elution profile, leading the authors to conclude that this mutant is unable to undergo the typical pH switch of HSP47 (18). In the study presented here, we do see that His\(^{215}\) and His\(^{216}\) are important for the general HSP47-collagen affinity, but we cannot detect any effect on the pH dependence of these residues; the pH dependence of the \(k_{\text{off}}\) and \(K_D\) of the H215N/H216N double mutant shows a similar behavior as the wild type, with \(k_{\text{off}}\) increasing from 0.094 s\(^{-1}\) at pH 7.5 to 0.273 s\(^{-1}\) at pH 6.0, but compared with the wild type, the binding is weakened by a factor of 2.3. We believe that the weaker binding explains the observed early elution from the collagen affinity column and does not imply that those residues play a role in the pH sensitivity of the interaction. For a mutant unable to undergo the pH switch, a delayed elution (i.e. at even lower pH) from a collagen affinity column would be expected.

If protonation of certain histidine residues triggers collagen release, removal of these protonation sites by mutagenesis should result in a dissociation rate constant (\(k_{\text{off}}\)) that is not (or at least is less) pH-dependent in comparison with the wild type. Only the double mutant H273N/H274N showed a decrease in the pH dependence of the \(k_{\text{off}}\) and the \(K_D\) values. This means that protonation at either site will decrease the affinity of HSP47 with collagen because both of the single mutants (H273N and H274N) show a similar pH-dependent behavior as the wild-type protein. In other words, protonation of either His\(^{273}\) or His\(^{274}\) is sufficient to induce collagen release. Interestingly, the double mutant H273A/H274A shows a behavior very similar to that of H273N/H274N, indicating that this is not a particular effect of the newly introduced asparagine side chains but rather of the missing imidazole moieties (i.e. the proton acceptor sites). However, in contrast to H273N, the single mutant H273A alone exhibits a small but significant reduction in pH dependence. This fits well with our observation from molecular dynamics analyses of wild-type HSP47, where His\(^{273}\) but not His\(^{274}\) was predicted to change its protonation state between ER and Golgi compartments, but opened the question of how His\(^{274}\) is able to compensate for His\(^{273}\) in the H273N mutant. Furthermore, His\(^{274}\) but not His\(^{273}\) is conserved in evolution.

Our computer simulations of the protonation states of His\(^{274}\) in the H273N protein indicate that the H273N mutation has a significant effect on the titration properties of His\(^{274}\). In the H273N mutant, His\(^{274}\) starts to be protonated at significantly higher pH values than in the wild-type protein (Fig. 10B), allowing His\(^{274}\) to compensate for His\(^{273}\) as a pH-dependent trigger. The finding that an asparagine is actually present at position 273 in several HSP47 orthologs supports the idea that this shift in the p\(_K_a\) of His\(^{274}\) might indeed be sufficient to allow His\(^{274}\) to exert a function in pH-dependent collagen release. In addition to an asparagine in birds and reptiles, in “fish” (clades Actinopterygii, Sarcopterygii, and Hyperoartia), tyrosine, leucine, and phenylalanine can be found at position 273. Our experiments using single amino acid substitutions in a canine protein background suggest that a tyrosine at position 273 can induce a shift in the p\(_K_a\) of His\(^{274}\) in a similar fashion as observed previously for H273N (Fig. 12), whereas phenylalanine and leucine cannot. For the H273F variant, this failure in pH-dependent release might easily be compensated for by the second ohnolog in spotted gar, which has the canonical tyrosine at position 273. Addi-
tionally, the phenylalanine carrying ohnolog has accumulated many additional amino acid exchanges. Compared with other fish sequences, the Tyr273 ohnolog is ~70% identical; the Phe273 variant, however, is only about 50% identical. This huge discrepancy might indicate that the Phe273 variant has lost its function as a collagen chaperon and might play a different role in the spotted gar. In cod and lamprey, there are no canonical ohnologs present, which could compensate for the lack of pH dependence of the H273L HSP47 variant. However, because pKa changes are highly dependent on the local environment, it still might be possible that Leu273 in the background of a fish protein is able to introduce the necessary pKa shifts to His274. These data suggest that only a limited set of amino acids is capable of conferring a suitable environment for His274 that allows this residue to act in pH-dependent collagen release in those species.

In this context, our earlier finding on the charged mutants at position 273 is also understandable (Fig. 11). Introduction of charged amino acids (H273D or H273K) at position 273 as well as significantly more hydrophobic side chains (Phe, Leu, and Ala) is less able to induce pH-triggered client release. In all of these cases, a reduced pH dependence is observed. This indicates that the H273A, H273K, and H273D mutations are unable to induce a shift of His274 pKa to higher values.

In other studies, conformational changes were often suspected of being important for the pH-dependent release. This was based on the differences observable in CD spectra under neutral and acidic conditions for recombinant mouse protein (15). However, we could only find minor differences in the overall CD spectra, thus indicating only minor structural changes over the pH range tested here (Fig. 9). We believe that the structural changes described previously (15) only occur at pH values below 6.0, an observation that has also been reported earlier (16). Most importantly, the less pH-dependent double mutant H273N/H274N shows a behavior in the CD spectrum similar to that of the wild type, hence excluding large structural rearrangements as the major driving force for pH-dependent client release. This is also confirmed by our steered molecular dynamics simulation (Fig. 8), in which dissociation of the HSP47 complex was easier for protonated HSP47 wild-type protein than for the non-protonatable H273N/H274N mutant. Here again, no major structural rearrangements were necessary to explain the effect.

Therefore, the mechanism of HSP47 dissociation in the Golgi apparatus seems to be driven by two factors. First, the protonation of HSP47 locally disrupts the collagen binding interface, most likely due to electrostatic repulsions. This leads to an increase in the koff and allows a rapid dissociation of the collagen client. In our experiments, we find a difference of ~7-fold in binding affinity between pH 7.0 and 6.0, translating into a small difference in the Gibbs free energy of about 5 kJ/mol. High resolution crystal structures will be needed to explain this in terms of an atomic model. Second, the released HSP47 is transported out of the Golgi in the ER via the KDEL receptor pathway, thus withdrawing the protein from the equilibrium. Our in-depth analysis of the protonation behavior of the HSP47 histidine residues and their influence on collagen binding and release identified positions 273 and 274 for the first part of this mechanism. Compared with the wild type, these two residues alone explain half of the increase in koff upon shifting the surrounding pH from 7 to 6. Most interestingly, His273 is not strictly conserved in all HSP47 orthologs; however, the residue at this position influences the pKa of His274.

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