Allometric scaling of skin thickness, elasticity, viscoelasticity to mass for micro-medical device translation: from mice, rats, rabbits, pigs to humans

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Emerging micro-scale medical devices are showing promise, whether in delivering drugs or extracting diagnostic biomarkers from skin. In progressing these devices through animal models towards clinical products, understanding the mechanical properties and skin tissue structure with which they interact will be important. Here, through measurement and analytical modelling, we advanced knowledge of these properties for commonly used laboratory animals and humans (~30 g to ~150 kg). We hypothesised that skin’s stiffness is a function of the thickness of its layers through allometric scaling, which could be estimated from knowing a species’ body mass. Results suggest that skin layer thicknesses are proportional to body mass with similar composition ratios, inter- and intra-species. Experimental trends showed elastic moduli increased with body mass, except for human skin. To interpret the relationship between species, we developed a simple analytical model for the bulk elastic moduli of skin, which correlated well with experimental data. Our model suggest that layer thicknesses may be a key driver of structural stiffness, as the skin layer constituents are physically and therefore mechanically similar between species. Our findings help advance the knowledge of mammalian skin mechanical properties, providing a route towards streamlined micro-device research and development onto clinical use.

With its highly accessible and abundant biological environment, the skin is an attractive site for a range of therapeutic applications for medical devices designed to improve healthcare. These include devices delivering drugs and vaccines into the skin1,2 and, alternatively, extracting skin biomarkers and electrical signals for diagnosis of disease3. In recent decades, the field has progressed beyond the needle and syringe towards more precise, minimally-invasive, micro-devices that exploit the mechanical properties of skin.

However, many existing and emerging micro-devices4,5 target the skin at scales and strain-rates distinct to the needle and syringe, exposing gaps in the knowledge of key mechanical properties. These knowledge gaps are further exacerbated as the path of medical device development progresses through typical animal models (e.g. mice, rats, rabbits and pigs) through to human clinical testing. To put simply, how do the key structural and mechanical properties of skin relate between different species? Identifying these properties would not only fill fundamental

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knowledge gaps, but could also streamline medical device research and development with reduced usage of animals and potentially smaller clinical trials.

Looking more closely at the challenge: it is well-established that skin is a complex, viscoelastic, biological composite structure, consisting of the epidermis (E), dermis (D) and hypodermis (H) or the subcutaneous tissue. The epidermis is further divided into stratum corneum (SC), a tough, physical and elastic barrier, comprised of keratinised dead skin cells, which makes accessing the layers beneath difficult. This is made even more challenging by the strain-rate dependent, viscoelastic viable epidermis (VE) – the uppermost living layer of the skin and dermis, where collagen, blood capillaries, antigen presenting cells (APCs) and biomarkers are located, often important for drug delivery and diagnostics. It is clear that the composition of skin is fundamental to its mechanical properties. However, because of strain-rate dependency, it is difficult to compare data directly with different experimentation protocols. Moreover, these skin layers differ in thickness between species, as shown in Table 1 (and a further table by Hirschberg et al.), where various measuring techniques and skin sites were used.

One particular example of emerging micro-devices susceptible to skin differences is the microneedle class of devices for drug/vaccine delivery and wearable diagnostics. The therapeutic attributes of many types of microneedles have been established in different animal models – including mice (e.g.10,13,14), rabbits (e.g.13,14), and pigs (e.g.15,16). However, how these therapeutic attributes are translated to widespread human testing (e.g.17,18) is reliant upon understanding how the key skin mechanical properties change between the species. Indeed, so far, microneedle utility in humans is limited, with extrapolated designs empirically derived for humans with varying degrees of success19. For example, Li et al. reported difficulty in penetrating through human skin, compared with rat skin, using the same application conditions20. A reason for this may be the lack of data to aid design translation to humans. Another reason may be that skin thicknesses vary greatly between the size of species, e.g. the SC is ~5 µm in mice to ~10–20 µm in humans, and also between sites within the same species20. This is despite almost all species possessing the same distinct epidermal, dermal and hypodermal layers3. A challenge in tissue mechanical property characterisation (such as elastic moduli) is that they often depend on test methods – orientation of loading conditions31, state of the sample (e.g. in vivo, ex vivo)32, testing rate34 and contact area35. As a result, for human skin, the reported elastic modulus can vary from 0.4–0.8 MPa for torsion in vivo36 to 4.6–20 MPa for tensile tests and 0.05–0.15 MPa for suction tests, according to Pailler-Mattei et al.35 Because of these factors, results from various skin studies are difficult to compare.

Nevertheless, within given species, the knowledge of relevant skin mechanical properties has been advanced – for example on elasticity37,38, fracture energy39 and percutaneous absorption40. However, in terms of mechanical properties of biological tissues, we are not aware of allometric investigations (the study of physiological characteristics relating to body size) of key mechanical properties of skin, ranging across body masses spanning orders of magnitudes (from ~30 g to ~150 kg), using a consistent approach. In this paper, we measured key skin physical and mechanical properties at relevant sites – thickness and elastic moduli at the microscale interface – using micro-indentation across five mammalian species of regular use in medical device research and development.

### Table 1. Selection of mean skin thicknesses of selected species reported in literature for non-weight bearing sites (mean ± SD/SE*).  

| Species | Site | SC | VE | D | Source |
|---------|------|----|----|---|--------|
| Mouse   | Dorsum | 9  | 29 | 662 | 48     |
| Mouse   | Buttck, ear, shoulder, back, abdomen (paraffin) | 3.38 ±0.30* | 11.50 ±1.24* | — | 48     |
| Mouse   | Buttck, ear, shoulder, back, abdomen (frozen) | 6.69 ±0.96* | 9.24 ±0.96* | — | 48     |
| Mouse   | Back | — | — | — | —       |
| Rat     | Dorsum | 18 | 32 | 2040 | 48     |
| Rat     | Buttck, ear, shoulder, back, abdomen (paraffin) | 4.04 ±0.47* | 15.34 ±1.21* | — | 48     |
| Rat     | Buttck, ear, shoulder, back, abdomen (frozen) | 9.91 ±1.14* | 10.70 ±1.73* | — | 74     |
| Rabbit  | Lumbar dorsum | 11.7 ±3.6 | 20.6 ±4.0 | 2174.0 ±486.7 | 74     |
| Rabbit  | Lumbar dorsum | 9.5 ±1.6 | 19.4 ±4.8 | 1719.3 ±258.5 | 74     |
| Rabbit  | Buttck, ear, shoulder, back, abdomen (paraffin) | 6.89 ±0.88* | 13.83 ±1.23* | — | 48     |
| Rabbit  | Buttck, ear, shoulder, back, abdomen (frozen) | 10.91 ±1.48* | 9.39 ±1.25* | — | 48     |
| Pig     | Buttck, ear, shoulder, back, abdomen (paraffin) | 12.85 ±1.19* | 53.17 ±3.19* | — | 48     |
| Pig     | Buttck, ear, shoulder, back, abdomen (frozen) | 41.33 ±3.73* | 15.37 ±1.51* | — | 48     |
| Pig     | Ear | 17–28 | 60–85 | 1440–2210 | (incl. H) 75 |
| Human   | Abdomen | 17 | 47 | 2906 | 48     |
| Human   | — | 10 | 50–120 | 2.28 | 76     |
| Human   | Various sites | — | 31–637 (incl. SC) | 521–1977 (E+D) | 41     |
Namely: mice, rats, rabbits, pigs (small and large) and humans (*in vivo* and *ex vivo*). Past indentation studies have shown the importance of accounting for tip-sample contact size for viscoelastic materials but have been used for the benefit of analysis simplification. Stress distribution on cartilage tissue was compared for flat and spherical tips with a cell death model and that while higher local stresses were experienced around the flat tip circumference, flat tips still provided comparable elastic moduli between the two tip profiles.

Our hypothesis is that skin of common laboratory animals – and humans – is mechanically a function of the layer thicknesses of each skin layer, and also related to body mass. To test this hypothesis, we first measured the SC, VE and dermis layer thicknesses. We then measured elasticity and viscoelasticity of full thickness skin. Finally, we aggregated these data to develop a simple model for estimating the elastic modulus of skin by knowing a basic parameter – the skin's layer thicknesses (relating to body mass). This relationship could be applied to more streamlined testing of mechanical micro-devices and other related work spanning from small to large animal models and indeed humans.

**Results**

To characterise the skin of different species, we first measured the thicknesses of the key skin layers – the SC, VE and dermis of mice, rats, rabbits, pigs and humans. We then indented the skin using a range of probes to measure the elastic moduli with consideration for viscoelastic effects, and used an analytical spring model to test the hypothesis that the elastic modulus is mainly dependent on skin thickness. We defined elasticity, modulus and material stiffness as the elastic modulus $E$, whereas the structural, axial or layer stiffness, dependent on the geometry of the material, $k$. The full skin thickness comprising the SC, VE and dermis was measured as a full thickness material interfacing medical devices, and analysed as a composite material with the three distinct layers. Values reported herein are mean ± standard deviation.

**Morphology.**  
Skin layers and thicknesses. Skin layer thicknesses measured using histology sectioning are shown in Fig. 1. Stained sections show structure, i.e. skin layers and cells residing within the skin strata, which were used as a guide to identify the skin layers. The SC is the top, stratified, corneocyte layer that is pink in colour visible in the 40x images Fig. 1(a–e iii). The VE is the layer beneath the SC in dark purple, which appears undulating at the lower boundary. Individual cells and nuclei (dark purple) can also be seen in higher magnifications. The dermis is the light purple region below the VE, which is the thickest of the three. Hair follicles/shafts (circular/oval white spaces in the dermis) can also be identified in all species apart from human skin.

Thickness measurement are collated in Table 2 and shown on a logarithmic scale vs. mass (Supplementary Table S1) in Fig. 1(f). While the three skin layers possess comparable thickness ratios between species (mean of $1:4.95$ of SC:VE:D), different features are observed between the species, which include:

- Rabbit dermis contains large numbers of hair follicles at $\sim4–5$ mm$^{-2}$, in contrast to pig dermis at $<1$ mm$^{-2}$.
- Hair follicle shafts of pig skin are larger than that of smaller species.
- Hair follicles were not observed on human skin sections.
- The rabbit skin features a relatively thick epidermal layer, closer to pigs than mice.
- Pigs of higher body mass ($\sim130–150$ kg) appear to have thicker VE than smaller pigs ($\sim20$ kg). The overall dermis thickness between both groups remain close ($P = 0.6806$).

Calculated $P$-values of layer thickness comparisons are provided in Supplementary Table S2. Notably, mice and rat skin are alike at $\sim20\mu m$ in the VE, as well as that of rabbits and small pigs despite the large differences in mass ($\sim20$ x and $\sim6$ x respectively). From the measurements of skin layer thicknesses, allometric scaling relationships fitted to experimental trends are presented as power law curves in Table 3(i) for skin thickness interpolation, if the mass of an animal is known.

**Skin mechanical properties.**  
Viscoelastic properties. Viscoelastic properties provide information on skin deformability at different strain rates and were required to obtain the elastic properties independent of the indentation rate. Figure 2(a) shows examples of the typical fit of Prony series to a subset group of raw data (human skin, indentation rate $v = 0.1$ mm s$^{-1}$, tip radius $R = 0.18$ mm). Prony series goodness of fit ($R^2$) are presented in Supplementary Table S3. Higher noise and spread can be seen in human skin experiments due to the lower than expected load measured (hence higher relative noise from the load cell and lower $R^2$). Small body movements and breathing from volunteers from *in vivo* human skin group were detectable and an example is shown in Fig. 2(b). Because of this, it was more difficult to fit curves to the *in vivo* tests and more optimisation had to be performed by adjusting the initial fitting parameters for the results to converge, which could suggest that the relaxation profiles are noticeably different to the rest of the species in Fig. 2(b). Raw force-time data of every replicate for each species is plotted in Supplementary Fig. S1.

Immediately after a step load application, we observed a transient, rapid drop of resistance force against the tip over the first second for all skin, followed by a steady-state, plateauing decrease for the remainder of the experiment. The first ten seconds (transient and some steady state phases) that are most applicable to skin-targeting devices are shown in Fig. 2(c,d). This behaviour was observed throughout the experiments for all skin types. The figure shows the force decreasing over ten seconds immediately after a $100\mu m$ s$^{-1}$ step load when the tip is held at constant (maximum) displacement of approximately 10% of the skin thickness. We observed that the skin of smaller species (e.g. mice, rats) show more viscoelastic effects than larger species (e.g. humans), i.e. the force decreases more over the same period (mean residual force after ten seconds is $\sim40\%$ for mouse skin but $\sim80\%$ for *in vivo* human skin). The *in vivo* human skin appeared not to relax as much compared to the other groups initially, but the at ten seconds it appears to continue decreasing more than the *ex vivo* human skin. Comparing between small and large pigs, the larger showed higher relaxation compared with the smaller. The Prony series parameters...
\( \tau_1, \tau_2, g_1, g_2 \) are produced in Table 4 and their resulting \( g(t) \) coefficient at 100 mm s\(^{-1} \) are given in Supplementary Table S4 for reference.

**Elastic properties.** The elastic moduli of skin were calculated as per Materials and methods - indentation procedure, accounting for the viscoelastic effects quantified previously. Figure 3(a) shows a typical fit of the Ogden model to raw data and Fig. 3(b) shows an *in vivo* human skin example. Greater data spread and noise were
**Table 2.** Measured skin layer thicknesses for comparison with literature values in presented Table 1.

| Species               | Site          | SC (µm) | VE (µm) | D (µm) | Approximate total (µm) |
|-----------------------|---------------|---------|---------|--------|------------------------|
| Mouse                 | Flank         | 4.19    | ±1.79   | 17.50  | ±4.98                  | 182.4 ± ±46.72 | 204 |
| Rat                   | Flank         | 9.38    | ±2.21   | 23.58  | ±9.79                  | 382.42 ± ±142.49 | 415 |
| Rabbit                | Flank         | 12.32   | ±3.52   | 84.34  | ±38.28                 | 1085.85 ± ±578.52 | 1183 |
| Pig (small)           | Ear           | 17.01   | ±3.96   | 89.60  | ±26.70                 | 1423.94 ± ±522.80 | 1531 |
| Pig (large)           | Ear           | 20.02   | ±3.55   | 131.50 | ±41.29                 | 1340.59 ± ±411.79 | 1492 |
| Human                 | Abdomen       | 17.07   | ±4.56   | 99.80  | ±49.29                 | 2284.05 ± ±1161.64 | 2401 |

**Table 3.** Parameters for fitted power law curves ($y = ax^b$) of measured skin layer thicknesses $y$ (µm) with respect to species mass $x$ (kg).

| Parameter                     | $a$   | $b$   | $R^2$ |
|-------------------------------|-------|-------|-------|
| (i) Skin layer (with large pigs) |       |       |       |
| Stratum corneum              | 10.01 | 0.143 | 0.96  |
| Viable epidermis             | 47.7  | 0.202 | 0.91  |
| Dermis                       | 756   | 0.187 | 0.71  |
| (ii) Skin layer (without large pigs) |       |       |       |
| Stratum corneum              | 9.98  | 0.147 | 0.97  |
| Viable epidermis             | 48.1  | 0.194 | 0.85  |
| Dermis                       | 617   | 0.310 | 0.98  |

captured during the indentation because of volunteer movements. Oscillations were likely caused by breathing, as observed that each inhalation corresponded to a slight rise of the load (and vice versa with exhalation and slight drop), increasing the difficulty in the Ogden fit. Despite this, both Prony and Ogden curves were still able to be fitted to the mean of the sinusoidal heart beat to obtain mechanical properties. Summaries of the goodness of fit and the $a$ coefficients are shown in Supplementary Tables S5 and S6 respectively. Raw force-displacement curves of every replicate for each species are shown in Supplementary Fig. S2.

Figure 3(c) shows mean and standard deviation force-displacement curves for all indents of all species. Curves followed behaviour typical of hyperelastic materials – linearly increasing for initial small displacements, then increasing more rapidly at large strains. Curves did not show a sudden drop in force during loading, which would have suggested puncturing of the skin. While all skin types followed the same loading path for the initial 0–5 µm indent, the human skin curves (both in vivo and ex vivo) had lower modulus compared with the other species. The maximum load at ~10% strain (maximum displacement) for human skin in vivo is 8.3 ± 5.7 mN and 2.24 ± 1.17 mN for human skin ex vivo, compared to 25.9 ± 33.3 mN for large pigs, 11.9 ± 12.2 mN for small pigs, 11.3 ± 12.5 mN for rabbit, 0.694 ± 0.348 mN for rat and 0.497 ± 0.164 mN for mouse.

In Fig. 3(e), we observed elastic modulus decreased with increasing tip radius in a log-log relationship. These trends were fitted as power curves in Table 5(i). Elastic modulus also increased with respect to species size for a given tip contact area, except for human skin being lower than rabbit skin. Large pig skin has the highest modulus of all tested species, whereas the skin’s modulus of small pigs followed more closely to other species. Such spread and variation of the elastic moduli were also reported in the indentation work of Ranamukhaarachchi et al.33, for example. Supplementary Fig. S5 shows the individual pig skin elastic moduli that pigs 4–5 had elastic moduli approximately one order of magnitude higher than pigs 1–3.

Table 6 shows that ex vivo human skin group had very low elastic moduli compared to other species in Fig. 3(c). To verify these were not anomalous samples, two additional ex vivo samples were repeated. A further group of in vivo human skin also was tested. In vivo human skin did not follow typical viscoelastic behaviour, i.e. slower tests gave higher measured forces, despite most other cases showed higher modulus with the faster indentation rate. The allometric scaling relationship to body mass is shown in Fig. 3(f), with curve fits presented in Table 5(iii).

**Linear analytical model.** To further help interpret the experimental trends, a simple model was used to help express the skin as three springs in series, isolating the elastic component. Skin properties of mice from Crichton et al.33 were used for each skin layer to estimate the elastic moduli and displacement of each animal and human model to test our hypothesis that the main influence of skin elasticity is the skin layer thicknesses.

Figure 4(a) shows spring model elastic moduli was in similar orders of magnitude compared with Ogden fitted experimental data, matching more closely towards smaller tip contact areas. This established a relationship where the species mass (which leads to skin thickness) could be used to determine the skin’s elastic modulus from experimental trends. Figure 4(b) shows that the amount of load taken up by the SC was approximately 65–90%, followed by the VE at 5–25% and the VE at 1–8%, where the modulus contribution to the full thickness skin can be seen. Increasing the tip radius reduced the modulus contribution by the dermis, but was the opposite for the
epidermal skin layers. Finally, the analytical model also showed that the dermis of larger species made less contribution to the overall modulus. For comparison Fig. 4(c) shows the analytical allometric scaling estimates for elastic modulus by body mass.

Discussion
The main purpose of this paper is to identify a way in which skin-interacting micro-devices can be translated easily between animal species to humans, and in the process, report on the physical, elastic and viscoelastic properties of skin across species. We hypothesised a relationship between species, so that their skin's thickness and mechanical properties could be estimated simply by knowing the body mass of that species through interpreting the data using a simplified analytical model.
Our results established – for the first time – experimentally-determined trends between these properties across four commonly used laboratory animal models to humans, spanning from micrometres to millimetres on skin thickness and body mass ranging from ~30 g to ~150 kg. We are not aware of such a broad study of this sort in present literature, on key properties with an allometry theme. Indeed, Wang et al. reported structural and mechanical properties (thickness, stiffness and modulus over body weight and skin sites) of only one species (mouse model) under compression. The concept of allometry was introduced over a century ago, and more specifically, allometric scaling models for organisms’ body mass and metabolic rates were proposed in recent decades. This concept was further discussed on the applicability of in vitro models by Ahluwalia. We propose to extend this relationship further to other physiological parameters in this study, namely, correlating skin’s thickness and its mechanical properties to body mass.

Figure 3. (a) Representative examples of raw data fitted to Ogden model. (b) Representative example of force-displacement curves when measuring human in vivo illustrating cyclical vibration caused by volunteers. (c) Mean force-displacement curves from all indents. (d) Mean force-displacement fitted to Ogden model normalised to 10% strain for all species with SD in shaded region. Note that y-axes have different scales. (e) Elastic moduli experimental trends shown vs. indentation tip radii with individual data points. (f) Elastic moduli shown vs. body mass.
In terms of variation in the spread of data observed (for example, in Fig. 3(d)) are commonly encountered with biological specimens\textsuperscript{36,37}, when compared to standard engineering materials (e.g. steel, rubber). However, substantial replicate variation, heartrate and respiration were also apparent in the indentation works of Miller et al.\textsuperscript{45} and Sridharan et al.\textsuperscript{38,39}, consistent with our results. Ranamukhaarachchi et al.\textsuperscript{27} also reported higher scatter in elastic modulus during indentation tests in porcine vs. human skin\textsuperscript{27}. We propose this could be due to environmental factors and skin layer thickness ratios. To boost data confidence, five specimens with five repeats per condition (indentation tip size and speed), with an additional two specimens for human ex vivo, were incorporated into the experimental design. Indeed, scatter of soft tissue data were also commented by Mattei and Ahluwalia\textsuperscript{40}, who discussed potential sources for variation that could affect measured mechanical properties. They also presented guidelines on minimising this effect, such as using animals from controlled environments (e.g. same breeding facilities), using same test methods and minimising sample preservation period were implemented in this study ensuring reproducible data that is applicable and comparable across the scientific community.

In terms of the indentation method and the pre-load using the Instron, except for human skin, the pre-load (~1 mN) was much smaller (~20x) the peak loads experienced during indentation. The pre-load was the smallest possible load used as a reference point to commence indentation and was unlikely to result in any significant changes to the data. While the pre-load was relatively large for the case of human skin, the properties of the skin was still measured and that the elastic modulus was the lowest of all skins tested using the Instron. A significant pre-load magnitude would however likely result in an increase in mechanical properties\textsuperscript{29}, preventing comparisons with other skins.

On the relation between skin thickness and body mass, we observed a proportional trend, as shown in Fig. 1(f). An exception is the dermis of the larger pigs is thinner than the smaller pigs, although the overall skin thickness was observed to be relatively close to each other. One explanation is that skin thickens up to a certain

| Parameter | $a$ | $b$ |
|-----------|-----|-----|
| (i) Body mass for various tip radii (measured) |  |  |
| 0.18 | 85704 | 0.1097 |
| 0.315 | 46881 | 0.1023 |
| 1 | 4375.2 | 0.1288 |
| 3.15 | 1380.4 | −0.1179 |
| (ii) Body mass for various tip radii (analytical model) |  |  |
| 0.001 | $8.2985 \times 10^6$ | 0.0221 |
| 0.01 | $1.7701 \times 10^6$ | 0.0284 |
| 0.1 | $3.6903 \times 10^5$ | 0.0342 |
| 0.18 | $1.9275 \times 10^5$ | 0.0359 |
| 0.315 | $1.2134 \times 10^5$ | 0.0359 |
| 1 | 45920 | 0.0359 |
| 3.15 | 17061 | 0.0359 |
| (iii) Tip radius for various species (measured) |  |  |
| Mouse | 672.98 | −1.900 |
| Rat | 2275.1 | −1.186 |
| Rabbit | 7943.3 | −1.441 |
| Pig (small) | 12618 | −1.227 |
| Pig (large) | 39355 | −1.152 |
| Human | 3311.3 | −1.584 |
| Human (in vivo) | 3499.5 | −2.002 |

Table 5. Parameters for fitted power law curves ($Y = ax^b$) of elastic moduli $Y$ (Pa) (i) measured and (ii) analytically modelled with respect to body mass $x$ (kg) or (iii) tip radii $x$ (mm).

| Tip radius (mm) | 0.18 | 0.315 | 1 | 3.15 |
|----------------|------|------|---|-----|
| Species | $E$ (kPa) | SD | $E$ (kPa) | SD | $E$ (kPa) | SD | $E$ (kPa) | SD |
| Mouse | 13.22 | 6.8 | 3.97 | 1.6 | 0.59 | 0.2 |  |
| Rat | 17.04 | 5.2 | 9.98 | 4.6 | 0.97 | 0.5 |  |
| Rabbit | 94.12 | 74.0 | 42.17 | 21.7 | 7.79 | 4.1 | 1.29 | 1.1 |
| Pig (small) | 102.52 | 47.2 | 55.09 | 38.9 | 9.70 | 6.4 | 0.43 | 0.4 |
| Pig (large) | 274.68 | 309.4 | 174.69 | 201.9 | 10.55 | 9.0 | 3.68 | 2.6 |
| Human | 50.26 | 19.3 | 19.83 | 7.1 | 4.74 | 1.9 | 1.99 | 1.1 |
| Human (in vivo) | 108.19 | 140.3 | 35.42 | 60.6 | 3.07 | 1.5 | 0.58 | 0.3 |

Table 6. Summary of mean elastic moduli for all species.
age or mass and because these pigs were raised for consumption (hence significantly increased body mass), their mass deviated away from their “normal” body mass over a relatively short period of time. For an experimental correlation of skin layer thicknesses relative to body mass, if we modify Table 3(i) to exclude larger pigs, we obtain a better-fitting allometric relationship (Table 3(ii)) plotted in Fig. 1(f).

Despite proposing this trend, it may not work well with species faraway from the boundaries of our chosen species, which could be a subject of further investigation. Nevertheless, all five species selected exhibited comparable ratios of skin layer thicknesses, i.e. mean of ~1:4:95 of SC:VE:D. This agrees with Lee and Hwang's reported ratios (3.7–16.8% of epidermis within the entire human skin for most regions)41.

In terms of factors affecting skin thickness: previous studies have shown that skin site, age42; gender20, BMI43 and SC hydration44 as possibilities, however, we propose that these differences are likely to be smaller than comparing across species (except weight-bearing extremities e.g. thicker SC around the palms and soles than the arms and abdomen)44. For instance, full-thickness human skin can decrease in thickness by ~300 µm between 20 to 90 years of age45, which is only up to ~10% difference of its full thickness. In this study, human skin thickness data exhibited the highest coefficient of variation of up to 50%; the highest among other species. This high inter-donor dependency was also observed in Mattei et al.’s study on liver tissue structure 46, who also commented that this was typical of human derived-samples. This could be minimised by using animals from the same sources, as with this paper (apart from human skin). While we compared different sites of animals, low intra-donor dependency was expected. This is supported by Lee et al. who showed that skin of the abdomen region is similar in thickness to the forearm (1332 ± 254 µm, vs. 1133 ± 215 µm respectively) and most other parts of the body41. We confirmed this in pigs to ensure this was the case with three sites on pig skin (data shown in Supplementary Fig. S6). The closest body of work we are aware of comparing tissues cross-species was by Malda et al. investigating the relationship between articular cartilage thickness and body mass47 and Monteiro-Riviere comparing the epidermal thicknesses of various species48.

On viscoelasticity, the skins of the smaller animals (i.e. mice, rats) were found to be more viscoelastic than those of larger species (Fig. 2(c)), which we hypothesised this was likely due to considerably higher relative thickness of the VE vs. dermis in smaller species (Table 2). Our relaxation profiles cf. Crichton et al.28, both showed a rapid decrease in normalised force to approximately 0.7 of the original force, then a slow decrease towards ~0.4 at 10 s and did not appear to be influenced by tip contact area. Furthermore, Crichton et al. showed that the VE is the most viscoelastic of all three skin layers33. Jee and Komvopoulos suggested time-dependent deformation of skin is mainly attributed by the cellular epidermis and dermis layers49. From our study, we suggest that the VE is the main contributor of viscoelasticity in full-thickness skin, as a more prominent viscoelastic behaviour is related to higher VE proportion in the skin composition (Table 2). An example would be the VE in full thickness skin of mice and rats were higher at ~4–9% and 6–9% for small and large pigs, whereas for rabbits and humans, this figure was lower at ~2–2.5%.
Furthermore, both the relaxation profiles of small and large pigs resembled each other and appeared closer to mouse and rat skin rather than to humans (as seen in the normalised load after 10 s in Fig. 2(c)). Despite differences in mass, we found relaxation profiles of rabbit skin behaved similar to human skin. We interpret these as likely due to fluid within the living layers of the skin i.e. intercellular space, and to a lesser extent the presence of collagen within the dermis, also known to behave viscoelastically. Age could be another factor, as while the VE of smaller pigs was lower than larger pigs, the relaxation curves appeared similar – younger skin may be more viscoelastic. Interestingly, Prony coefficients did not appear to vary much between species (0.65–0.78), suggesting allometric scaling trends between species may not be prominent.

On elastic modulus, we anticipated the skin’s elastic modulus would increase with species mass, however, human skin was one of the most compliant. We propose that variations in skin layer thickness ratios of humans affected this (i.e. greater VE layer influence, which was found to be the most compliant of all three layers), although other factors such as hair follicles, captive or environmental exposure differences and skin sites (to a lesser extent) may be relevant. Skin deformation contributed by the SC primarily was suggested by Jee and Komvopoulos. Indeed, this agreed with Fig. 4(b) that the SC was the main layer for stiffness contribution, and that thicker SC corresponded to higher elastic modulus (e.g. small vs. large pigs). Comparing our data with literature, Crichton et al. reported 860 kPa for mouse skin, using Equation 4, which was the trend fitted to the experimental data, we obtained a slightly lower estimate of 53 kPa. However, another example reported pig skin reduced modulus of 3.77 MPa (tip radius 20 μm), matching well with our estimate of 2.52 MPa (small pigs) and 5.69 MPa (large pigs). Overall, we have shown that the composition of skin is an important factor in influencing the mechanical properties of full thickness skin. Regarding allometric scaling of elastic modulus to body mass presented in Fig. 3(f), the smallest three tips showed consistent logarithmic increase, while the largest tip exhibited a decrease, although this was most likely due to limitations of testing very large tips on mouse and rat skins (Materials and methods – indentation tips).

In terms of scale effects, we found increasing tip contact area correlated with decreasing tissue elastic moduli shown in Fig. 3(e). This tip to modulus trend plotted as log-log gradients for each species showed similar slopes. Previous studies showing this trend proposed that corneocytes of the SC were distributing loads more effectively at smaller contact areas due to corneocyte-to-tip scales. Smaller pigs were found to have lower elastic modulus than larger pigs, despite similar overall skin thickness, suggesting scale effects could be more pronounced with certain skin layers, cell structure at different ages, or simply due to higher biological variation in large pig skin specimens, as shown in Supplementary Fig. S5 (with differences between specimens at approximately 1–1.5 orders of magnitude).

On the analytical model (Results - linear analytical model), it provided fresh insights into the skin’s elastic modulus as a function of animal mass across the species tested. For example, by knowing the elastic modulus of a mouse, we could approximate for humans with the mass. The purpose to develop the model was to help interpret our results; to understand the contribution of tissue layers, rather than to provide accurate solutions to solve for the skin properties (for this approach, alternative methods such as finite element analysis would be employed). As such, this model was purely elastic and did not account for viscoelastic and other effects, but still gave reasonable correlation between species. By normalising the data of tip radius and skin thickness, a clearer relationship between elastic modulus and species could be observed, supporting our hypothesis that the main driver of skin’s elastic modulus is the thickness of the skin layers. Other more complex skin modelling approaches utilise finite element analysis (e.g. which is time consuming and expensive. For many purposes, this model serves as an engineering tool that can provide estimates of the skin’s mechanical properties. Limitations to this model may include inapplicability of species from harsher habitats or outside the scales herein. For example, an elephant weighing ~4 tons, has a skin thickness of ~17.5 mm, but Equations 5–7 suggests a full skin thickness of approximately half at 9.7 mm. However, other species within the mass range investigated, such as sheep, which typically weighs ~40 kg at adulthood, has a skin thickness of 1.83–2.15 mm. This compares well with the same formulae, which estimated a full skin thickness at ~2.3 mm. Based on this information, if we use our estimated mean SC:VE:D ratio of 1:4:95, using Equations 15–17, we could estimate that indenting using a tip radius of 0.18 mm would give a bulk elastic modulus of sheep skin of 422 kPa, which is comparable to Manan & Mahmud’s study who reported a tensile modulus of 369–539 kPa.

In terms of freshness of skin and its mechanics, we endeavoured to work with freshly excised skin. However, ex vivo human skin samples obtained were not able to be tested immediately after surgical excision and were refrigerated for up to 24 hours before indentation. Some studies show differences in tissue mechanical properties after storage in refrigerated conditions, and changes to Young’s modulus after storage in freezing conditions. However, studies such as Banga showed minimal changes to tissue quality stored under refrigeration that were used within a few days after excision. Regardless, we did not observe any visual degradation to sample quality between obtaining the skin and completing the experiment, and that current protocols obtaining and processing skin are unlikely to have influenced tissue mechanics. Further, to demonstrate data applicability to in vivo human conditions, we compared ex vivo skin with in vivo conditions with volunteers. In force-relaxation experiments, in vivo human skin showed less decrease in force over the initial 2–3 seconds vs. ex vivo conditions. We hypothesised additional pressure from blood circulation and skin in its original tension and state could contribute to this difference. Previous porcine studies on brain tissue also showed discrepancies in material stiffness between tissue states. This is particularly relevant where development of new devices is performed on excised tissue, with the expectation that translation to live humans will be simple. Despite this, we propose they are unlikely to affect most tests of micro-scale devices, as the differences are small; ex vivo skin could still provide valuable data when emulating in vivo conditions for certain purposes, such as mechanical testing.

Mechanical testing of medical devices often commences on small animal models during pre-clinical stages. Our data can help enable rapid translation from laboratory conditions by streamlining and reducing animal usage before human testing with improvements being made to devices more effectively. Based on date presented here,
we highlight that \textit{ex vivo} human skin, in combination with rabbit and/or small pig skins may be suitable for testing micro-medical devices in developmental stages to obtain a translatable elastic and viscoelastic result. Indeed, the ability to relate animal model experiments to human clinical applications is applicable to several medical interventions, such as in surgery and soft tissue cutting, impact and puncture of skin, and skin based diagnostic and sensing devices, which can be optimised to target the skin at a more precise and effective level.

Conclusions
To address the need for effective and rapid translation of micro-scale medical devices from laboratory to clinics, we investigated the morphology and the mechanical properties of the skin of mice, rats, rabbits, pigs and humans; with histology and indentation. We hypothesised that we could identify a relationship between these species for elastic modulus and skin thickness using a simple allometric correlation to species mass. We reported this relationship using experimental trends, which correlated well with reported literature. Our data suggest that the thickness of each layer of skin increased with species mass, together with the elastic modulus (except for human skin). We extended this to humans and observed a reduction in material stiffness from \textit{ex vivo} skin to \textit{in vivo} skin. Using indentation, our measured elastic moduli of the selected species is between $10^2$–$10^6$ Pa with indenter tip radii between 0.18–3.15 mm. From our study, we recommend that \textit{ex vivo} human skin, rabbit skin and small pig skin would be suitable for pre-clinical testing of medical devices. By establishing the relationships between animal models and humans, we can help translate devices more rapidly through the knowledge of understanding the mechanical properties of the skin and its reaction to a micro-scale like device acting on its surface.

Materials and Methods
In this paper, we used indentation to measure elastic and viscoelastic properties of skin specimens from mice, rats, rabbits, pigs and humans. We then fitted the Ogden hyperelastic model and two-term Prony series to the loading and force-relaxation curves, to obtain these properties respectively.

Skin tissue preparation for indentation. Skin was collected from five species: mouse (flank), rat (flank), rabbit (flank), pig (ear) and human (abdomen) spanning four orders of magnitude in mass: mouse ~30 g, rat ~300 g, rabbit ~3 kg, pig ~30 kg, and humans at ~70 kg. In addition, larger, abattoir sourced pigs ~130–150 kg were compared (i.e. from animal experiments and commercial abattoir). This provided comparison between the same species but at two body masses. The mass of the first four species were measured directly from the animals (large pigs were quoted directly from the abattoir), and humans for \textit{ex vivo} skin were estimated from Walpole \textit{et al.}.\textsuperscript{59}

The mass of the human (\textit{in vivo}) was collected directly from volunteers.

Mice (CD1, female, 10 ± 1 weeks old), rats (Wistar, female, 12 ± 1 weeks old), rabbits (New Zealand white, female, 12 ± 2 weeks old) and small pigs (~20 kg) (Large White, female, 9 ± 1 week old) were obtained from the University of Queensland Biological Resources (St Lucia QLD, Australia). Skin sites selected were from large, uniform areas of the body and avoided weight-bearing regions with thicker SC thicknesses.\textsuperscript{41} Flank (mouse, rat, rabbit) or dorsal ear (pig) skin tissue was excised for testing immediately post euthanasia (mouse/rat with CO\textsubscript{2} chamber, rabbits and pigs with overdose of ketamine/xylazine). Ear skin of large pigs (Large White female, >1 year old) was purchased from Highchester Meats Ltd (Gleneagle QLD, Australia) with skin excised from dorsal ear cartilage without post-cull treatment i.e. hot water dip. Human skin was sourced from the Princess Alexander Hospital (Herston QLD, Australia) from female abdominoplasty patients aged 36 ± 7.8 years old (mean ± SD). Animal hair was removed with hair clippers (Pet grooming kit, Wahl, Stirling IL, USA) followed by a razor blade shave (Xtreme3, Schick, St Louis MO, USA). Fat was removed from the skin by scalpel. \textit{In vivo} volunteer human skin (dorsal forearm with no visible scarring or defects) was also compared against \textit{ex vivo} human skin (healthy 3 males and 2 females, 24 ± 1.5 years old, mean body mass 63 ± 7.6 kg).

Mechanical testing of skin was collected within three hours post euthanasia, except for pig and \textit{ex vivo} human skin, where supplies were not available on demand – testing was completed within 48 hours of obtaining skin samples. In this situation, skin samples were excised with hydration and viability maintained, similar to Jee and Komvopoulos\textsuperscript{37}, except placing the bottom side of skin on cell culture media (RPMI 1640 Medium, Gibco, Thermo Fisher Scientific, Waltham MA, USA) (not submerged) with antibiotics (Ampicillin, Gibco, Thermo Fisher Scientific, Waltham MA, USA) refrigerated at 4 °C. It was ensured the surface was dry to avoid potential changes to the epidermal mechanical properties\textsuperscript{38,40}. Skin was returned to room temperature before testing.

All animal work carried out has been approved by the University of Queensland Animal Ethics Committee (ethics number ANRFA/AIBN/473/15). All human work carried out has been approved by the University of Queensland Human Research Ethics Committee (ethics numbers 2008001342 and 2017000693). Written informed consent was obtained from all participants. All experiments were carried out in accordance with the University of Queensland guidelines and regulations.

Histology. Five individual skin samples of each species were collected for skin thickness measurement. Frozen-section method was selected over paraffin due to less exposure to processing and faster turnaround time. Subcutaneous layer was removed during the dissection. Skin was cut to ~1 cm\textsuperscript{2} size and submerged in 10% neutral buffered formalin (NBF) (HT501128, Sigma Aldrich, St Louis MI, USA) following standard histology protocol\textsuperscript{19} immediately after harvest. Samples were embedded in moulds (Peel-A-Way, Polysciences, Warrington PA, USA) with sectioning matrix (Tissue-Tek OCT, Sakura Finetek, Alphen aan den Rijn, the Netherlands) and frozen by liquid nitrogen. Samples were pinned during fixation and held up right during freezing to ensure perpendicular sections were obtained at 14 µm thick (Microm HM 560, Thermo Fisher Scientific, Waltham MA, USA) and at least three slides were collected from each specimen (subject to quality of sections obtained). Between each slide (Superfrost Plus, Thermo Fisher Scientific, Waltham MA, USA), at least 350 µm of specimen was discarded so sections collected were not adjacent to each other.
Sections were imaged with confocal microscopy (LSM 510 META, Zeiss, Oberkochen, Germany) using 10x and 20x objectives and plain, white light to observe morphology and skin layers. An 800 nm laser was used to identify the presence of collagen at 430 nm, which indicates the approximate dermal layer 62, if layers were difficult to distinguish. Representative images are shown in Supplementary Fig. S7. Five replicates of each species were measured at least 20 times, up to 100 times on each skin layer depending on species and sample quality (i.e. no folding, curling and/or shattering of histology specimens) (Zen Black Edition 2009, Zeiss, Oberkochen, Germany). Distance was taken as perpendicular to the SC surface and spaced approximately three times the SC length apart between each measurement illustrated in Supplementary Fig. S8. Bright-field microscopy with haematoxylin and eosin following staining 63 was also used to identify the separate skin strata (BX45, Olympus Corporation, Tokyo, Japan), at 4x, 10x and 40x magnification. Brightness, contrast and colour balance were adjusted (Photoshop CC, Adobe Systems Incorporated, San Jose CA, USA).

Indentation equipment. Species were separated into thin and thick skin. For thin skin (mice and rats), a Triboindenter (Hysitron T900, Minneapolis MN, USA) with a MultiRange NanoProbe transducer was used. For the remaining, thick skin, a universal testing machine (Instron 5543, Norwood MA, USA) with a 5 N load cell were used. Both equipment has an overlapping indentation rate of 100 µm s$^{-1}$. The reason for the separation is due to Triboindenter’s maximum vertical displacement of ~90 µm, not deep enough for thicker skin. Secondly, for small displacements (e.g. up to 50 µm), Instron data contained relatively high levels of noise. For these indentations, the Triboindenter was used.

Polydimethylsiloxane (PDMS) as a handling layer for indentation. A PDMS layer was used as handling layer for the skin to be pinned on, and to protect the load cell in case of overshooting the intended displacement. PDMS backing was made using Sylgard 184 Silicone Elastomer Kit (Dow Corning, Midland MI, USA) mixed with the supplied curing agent in a 20:1 ratio. Vacuum chamber removed air bubbles in mixture. Mixture was poured in circular mould to 7–10 mm thick and cured in 60 °C oven for two hours.

Indentation tips. Custom-made, aluminium, flat cylindrical tips with radii of 0.180, 0.315, 1.000 and 3.150 mm were used. These radius sizes gave contact area one order of magnitude larger than the previous (except the 0.18 mm tip, which was the smallest we manufactured) to provide a range of readings and extrapolation of material properties outside the scales tested down to the sub-cellular scale 64. Tip sizes were selected based on Wayes et al., who indicated a suitable range of indentation tip size between 0 and 100% of the specimen thickness, as tip sizes significantly larger than the skin thickness changes the experiment to a flat plate compression model 65. Smaller tip sizes in the micrometre range are more closely associated to and more relevant to the typical scales of microneedle devices.

Surface roughness. The gap between skin and tissue due to surface roughness may affect mechanical analysis, however, to minimise this effect, skin furrow and hair root regions were avoided. Skin roughness surface amplitudes shown in literature were smaller than our indentation depths (e.g. mice $R_{a}$ (arithmetic mean) ~7.8 66, human $R_{p}$ (root mean) ~22–30 67).

Indentation procedure. A diagram illustrating the method is shown in Supplementary Fig. S9. Skin was placed on wet paper towel moistened with 1x phosphate buffered saline (PBS) during experiment to prevent dehydration 68. Mice and rat skin (with 1x PBS paper towel layer) were placed on a Triboindenter specimen holder. Rabbit, pig and human skin were pinned on the edges together with 1x PBS paper towel on a PDMS handling layer using hypodermic needles (to its original dimensions prior to excision to mimic in vivo conditions) for the Instron stage. Indentation area excluded regions near sample boundaries and the pins. Mice and rat skin were not pinned due to loose-skin nature of animals and limited working space in the Triboindenter. A mass-balance experiment (Supplementary Table S9) was carried out to verify skin was not overhydrated through passive capillary diffusion or osmosis. For the in vivo human skin experiment, the volunteers rested their arms on the Intron stage. Indentation depth was set at approximately 10% of the material thickness 66,67 to eliminate potential substrate effects while maintaining the ability to measure the effects of the full skin thickness. Tests performed using the Instron incorporated a preload of ~1 mN to ensure full contact of the tip and skin surfaces prior to commencing the loading ramp, similar to the automatic contact detection of the Triboindenter. Loading ramp was 0.01 mm s$^{-1}$ and repeated at 0.1 mm s$^{-1}$. This was not pre-conditioning the material, and the magnitude of the pre-load was the minimum readout from the load cell without ambient fluctuations. This was followed by a fixed-displacement hold for 10 s and an unload ramp at the same rate. We doubled the recording hold time (i.e. data points) for in vivo human skin to allow better curve-fit due to small body movements detected during measurement (Fig. 3(b)). Increasing the sampling rate allowed a reduction in recording/relaxation duration. Quality of fit ($R^{2}$) was also used as an indicator to determine the minimum required duration without reaching a fully-relaxed plateau. to determine Prony coefficients; force-relaxation could be as short as less than one second 68. Each condition was repeated five times for each tip size and indentation rate, with five replicates per species to ensure robustness of data allowing for natural variations in biological specimens.

Data analysis. Derivations to the Prony series and Ogden curve fits are from Crichton et al. and Lin et al.33,69 Force, displacement and time data were obtained from indentation. Two-term Prony series curve was fitted to force-time data of the hold section during indentation as per Wu et al.70, with the Prony series in general form (Equation 1) and in two-term form (Equation 2):
\[
G(t) = 1 - \sum_{i=1}^{N} g_i \left( 1 - e^{-\frac{t}{\tau_i}} \right)
\]  
(1)

\[
G(t) = 1 - g_1 \left( 1 - e^{-\frac{t}{\tau_1}} \right) - g_2 \left( 1 - e^{-\frac{t}{\tau_2}} \right)
\]  
(2)

in which \( g_i, \tau_i \) are relaxation magnitudes, \( \tau_1, \tau_2 \) are time constants, \( t = \frac{x_{\text{max}}}{v} \), where \( x_{\text{max}} \) is the maximum indentation displacement and \( v \) is the indentation velocity. Equation 2 gives a value between 0 and 1, which is used to obtain the reduced elastic modulus by multiplying the instantaneous elastic modulus with \( G(t) \). Replicates that did not converge for the code to filter ambient vibrations were discarded.

The Ogden hyperelastic model in Equation 3 was used to fit the force-displacement curves previously demonstrated by Lin et al.\(^69\) beyond small-strain definition, applicable for non-linear stress-strain behaviour of soft material indentations (although not dramatic, our range we indent to approximately 10% of the skin thickness). The authors also considered the Ogden model as the most appropriate for biological tissues\(^71\). Specifically, we fitted the model to the loading curve instead of the unloading curve for conventional indentation\(^66\), effectively characterising the material properties at the instance the skin is being loaded, in the same manner as a micro-scale medical device being applied onto the skin. Replicates unable to be fitted despite changing the ‘initial guess’ parameters (i.e. \( E \) and \( \alpha \)) and boundary conditions were discarded.

\[
P = \frac{40E_0a^2}{9\epsilon(1 - \nu^2)} \left[ \left( 1 - \epsilon^g \right)^{\alpha^2} - (1 - \epsilon^g)^{\alpha - 1} \right]
\]  
(3)

where \( P \) is the load, \( E_0 \) is the elastic modulus, \( a \) is the tip contact radius, \( \alpha \) is the fitting parameter, \( \nu \) is the Poisson’s ratio and \( \epsilon \) is the strain (instantaneous indentation depth/skin thickness) for large deformations applicable to this study\(^69\).

Matlab 2015a and 2016a (MathWorks, Natick MA) were used to automate data processing. Ogden and Prony curves were fit using the nlinfit function. Up to 10% of the initial force-displacement data was excluded from the origin to avoid fitting over noise/movement (in vivo) with relatively low forces, ambient noise and transient artefacts.

**Statistical analysis and graph plotting.** Skin thickness power law equations were determined using the Matlab (2016a, MathWorks, Natick MA) Curve Fitting Tool. Prism (GraphPad Inc., La Jolla CA) was as follows: (a) Plotting of all graphs. (b) Statistical significance in indentation and thickness data between species using ordinary one-way ANOVA multiple comparisons (Tukey’s multiple comparisons test). Statistical significance levels shown in figures and tables are: ns (\( P > 0.05 \)), * (\( P \leq 0.05 \)), ** (\( P \leq 0.01 \)), *** (\( P \leq 0.001 \)), **** (\( P \leq 0.0001 \)). Standard deviation is stated unless otherwise specified. (c) Elastic modulus power law equations were determined using the Nonlinear Regression (log-log line) Tool.

Power curves were fitted to the central trend of each species to obtain an allometric scaling relationship for skin thickness, elastic modulus and indenter tip radius from Tables 3 and 5:

\[
y = ax^b
\]  
(4)

\[
\log y = \log a + b \log x
\]  
(5)

in both power law and logarithmic forms, with the latter resembling the linear algebraic equation \( y = mx + c \) for the log-log graphs shown in Figs 1(f) and 3(e,f).

**Analytical model.** To investigate whether elasticity can be primarily defined as a function of skin layer thicknesses, the skin was modelled as three balanced, ideal springs in series, with no mass, damping or viscoelastic effects representing each of the skin layer and to isolate the system to the elastic components. Viscoelasticity could be applied later using the Prony coefficients as they do not appear to depend on scale\(^33\); including damping elements in the model but this complicates the calculation and deviates the model away from its intended purpose. The three spring system was comparable to the work of Pailler-Mattei et al., who quantified their tissue layers as dermis, hypodermis and muscle for their indentation study\(^68\). Starting from a basic spring force-displacement relationship with Hooke’s law:

\[
F = kx
\]  
(6)

A constant contact area throughout the skin equal to the tip contact area was assumed, i.e. 1D treatment of compression in this model. This was expressed as the material stiffness of an infinite series of springs in series fixed on one end\(^32\):

\[
k = \left[ \frac{1}{\sum_{i=1}^{n} k_i} \right]^{-1}
\]  
(7)

For a three-layer composite structure model defined as the SC, VE and D layers, this becomes:
\[ F = \left[ \frac{1}{k_{SC}} + \frac{1}{k_{VE}} + \frac{1}{k_D} \right] x \]  

where \( x \) is the displacement of the tip and \( k_{layer} \) is the stiffness of each layer as the axial stiffness in relation to elasticity:

\[ k_{layer} = \frac{E_{layer}A}{t_{layer}} \]

With \( A \) approximated as the tip surface area and \( t_{layer} \) as the measured thickness of the skin layer and \( E_{layer} \) as the elastic moduli of the SC, VE and D of mice skin obtained from mice by Crichton et al.33 fitted using power law (parameters in Supplementary Table S8). Mouse skin layer data was used here, as we hypothesised that skin of different species share common material properties.

A generalised equation to estimate the elastic modulus, if the skin layer thicknesses were known, was developed from the definition of stress, assuming an elastic relationship:

\[ E = \frac{\sigma}{\varepsilon} \]

\[ \sigma = \frac{F}{A} \]

Substituting Equations 8–11, we obtained the generalised analytical equation to estimate elastic modulus of skin:

\[ E = \frac{1}{\frac{1}{k_{SC}} + \frac{1}{k_{VE}} + \frac{1}{k_D}} A \varepsilon \]

where \( \varepsilon \) is the maximum indentation displacement, \( A = \pi R^2 \) for a flat, cylindrical tip and \( \varepsilon \) is the maximum strain. The area modifier allowed for tip interface scale dependencies as observed in biological tissues.

A schematic diagram of the simplified skin model is shown in Supplementary Fig. S11.

**Data availability**

The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

**References**

1. Kendall, M. A. The delivery of particulate vaccines and drugs to human skin with a practical, hand-held shock tube-based system. Shock Waves 12, 23–30 (2002).
2. Kendall, M. A. F. Needle-free vaccine injection. Handb Exp Pharmacol 197, 193–219 (2010).
3. Corrie, S. R., Coffey, J. W., Islam, J., Markley, K. A. & Kendall, M. A. F. Blood, sweat, and tears: developing clinically relevant protein biosensors for integrated body fluid analysis. Analyst 140, 4350–4364 (2015).
4. Cheung, K. & Das, D. B. Microneedles for drug delivery: trends and progress. Drug Deliv 1–17 (2016). https://doi.org/10.3109/10717544.2014.986309.
5. Larrateta, E., McCrudden, M. T. C., Courtenay, A. J. & Donnelly, R. F. Microneedles: A New Frontier in Nanomedicine Delivery. Pharm Res 33, 1055–1073 (2016).
6. Riema Pierre, M. & Rossetti, F. Microneedle-Based Drug Delivery Systems for Transdermal Route. Curr Drug Targets 15, 281–291 (2014).
7. Crichton, M. L. & Kendall, M. A. F. In Micro and Nanotechnology in Vaccine Development 343–356 (Elsevier, 2017). https://doi.org/10.1016/B978-0-323-39981-4.00018-X.
8. Hirschberg, H. J. B., van Riet, E., Bouwstra, J. A. & Kersten, G. F. A. Animal models for cutaneous vaccine delivery. Handb Exp Pharmacol 71, 112–122 (2015).
9. Fernando, G. J. P. et al. Potent Immunity to Low Doses of Influenza Vaccine by Probabilistic Guided Micro-Targeted Skin Delivery in a Mouse Model. PLoS ONE 5, e10266 (2010).
10. Coffey, J. W., Corrie, S. R. & Kendall, M. A. F. Early circulating biomarker detection using a wearable microprojection array skin patch. Biomaterials 34, 9572–9583 (2013).
11. Muller, D. A. et al. Inactivated poliovirus type 2 vaccine delivered to rat skin via high density microprojection array elicits potent neutralising antibody responses. Sci. Rep 6, 22094 (2016).
12. Li, H. et al. Microneedle-Mediated Delivery of Copper Peptide Through Skin. Pharm Res (2015). https://doi.org/10.1007/s11095-015-1652-x.
13. Chen, K. et al. Fabrication of Micro-Needle Electrodes for Bio-Signal Recording by a Magnetization-Induced Self-Assembly Method. Sensors (Basel) 16, 1533 (2016).
14. Kim, Y. C., Grossniklaus, H. E., Edelhauser, H. F. & Prausnitz, M. R. Intrastromal Delivery of Bevacizumab Using Microneedles to Treat Corneal Neovascularization. Invest Ophth Vis Sci 55, 7376–7386 (2014).
15. Lahiji, S. F., Dangol, M. & Jung, H. A patchless dissolving microneedle delivery system enabling rapid and efficient transdermal drug delivery. Sci. Rep 5, 7914 (2015).
16. Nguyen, J., Ra, K., Morra, M. & Popova, I. The Influence of Solid Microneedles on the Transdermal Delivery of Selected Antiepileptic Drugs. Pharmaceutics 8, 33 (2016).
17. Pearton, M. et al. Microneedle delivery of plasmid DNA to living human skin: Formulation coating, skin insertion and gene expression. J Control Release 160, 561–569 (2012).
18. Mohammed, Y. H. et al. Microneedle Enhanced Delivery of Cosmeceutically Relevant Peptides in Human Skin. *PLoS ONE* 9, e101956 (2014).
19. Kim, Y.-C., Park, J.-H. & Prausnitz, M. R. Microneedles for drug and vaccine delivery. *Adv Drug Deliv Rev* 64, 1547–1568 (2012).
20. Sandby-Moller, J., Poulsen, T. & Wulf, H. C. Epidermal Thickness at Different Body Sites: Relationship to Age, Gender, Pigmentation, Blood Content, Skin Type and Smoking Habits. *Acta Derm-Venereol* 83, 410–413 (2003).
21. Rittig, L. Cellular mechanisms of skin repair in humans and other mammals. *J Cell Commun. Signal.* 10, 103–120 (2016).
22. Choi, A. & Zheng, Y. P. Estimation of Young’s modulus and Poisson’s ratio of soft tissue from indentation using two different-sized indentors: finite element analysis of the finite deformation. *Med Biol Eng Comput.* 43, 258–264 (2005).
23. Urbanczyk, C. A., Palmeri, M. L. & Bass, C. R. Material characterization of in vivo and in vitro porcine brain using shear wave elastography. *Ultrasound Med Biol* 41, 713–723 (2015).
24. Briscoe, B. J., Fiori, L. & Pelillo, E. Nano-indentation of polymeric surfaces. *J. Phys. D: Appl. Phys.* 31, 2395–2405 (1999).
25. Chen, J. Nanobiomechanics of living cells: a review. *Interface Focus* 4, 20130055–20130055 (2014).
26. Pailler-Mattei, C., Bec, S. & Zhaozouari, H. In vivo measurements of the elastic mechanical properties of human skin by indentation tests. *Med Eng Phys* 30, 599–606 (2008).
27. Ranamukhaarachchi, S. A. et al. A micromechanical comparison of human and porcine skin before and after preservation by freezing for medical device development. *Sci. Rep.* 6, 32074 (2016).
28. Crichton, M. L., Chen, X., Huang, H. & Kendall, M. A. F. Elastic modulus and viscoelastic properties of full thickness skin characterised at micro scales. *Biomaterials* 34, 2087–2097 (2013).
29. Nicolopoulou, C. S., Giannoudis, P. V., Glaros, K. D. & BARBENEL, J. C. In vitro study of the failure of skin surface after influence of hydration and preconditioning. *Arch Dermatol Res* 290, 638–640 (1998).
30. Schnook, E. P., Meingassner, J. G. & Billich, A. Comparison of human skin or epidermis models with human and animal skin in vitro: percutaneous absorption. *Int J Pharm* 215, 51–56 (2001).
31. Tang, B. & Ngan, A. H. W. Accurate measurement of tip–sample contact size during nanoindentation of viscoelastic materials. *J. Mater. Res.* 18, 1141–1148 (2003).
32. Bae, W. C., Schumacher, B. L. & Sah, R. L. Indentation probing of human articular cartilage: Effect on chondrocyte viability. *Osteoarthr. Cartil.* 15, 9–18 (2007).
33. Crichton, M. L. et al. The viscoelastic, hyperelastic and scale dependent behaviour of freshly excised individual skin layers. *Biomaterials* 32, 4670–4681 (2011).
34. Wang, Y., Marshall, K. L., Baba, Y., Gerling, G. J. & Lumpkin, E. A. Hyperelastic Material Properties of Mouse Skin under Compression. *PLoS ONE* 8, e74399 (2013).
35. Alhuwalia, A. Allometric scaling in-vitro. *Sci. Rep.* 7, 42113 (2017).
36. Park, S. H. et al. Microneedle-based minimally-invasive measurement of puncture resistance and fracture toughness of sclera. *Acta Biomaterialia* 44, 286–294 (2016).
37. Jee, T. & Konvopoupolos, K. In vitro measurement of the mechanical properties of skin by nano/microindentation methods. *J Biomech* 47, 1186–1192 (2014).
38. Miller, K., Chinzei, K., Orssengo, G. & Bednarz, P. Mechanical properties of brain tissue in vivo: experiment and computer simulation. *J Biomech* (2000).
39. Sridharan, A., Rajan, S. D. & Muthuswamy, J. Long-term changes in the material properties of brain tissue at the implant–tissue interface. *J. Neural Eng.* 10, 066001 (2013).
40. Mattei, G. & Alhuwalia, A. Sample, testing and analysis variables affecting liver mechanical properties: A review. *Acta Biomaterialia* 45, 60–71 (2016).
41. Lee, Y. & Hwang, K. Skin thickness of Korean adults. *Surg Radiol Anat* 24, 183–189 (2002).
42. Branchet, M. C., Boisin, S., Frances, C. & Robert, A. M. Skin Thickness Changes in Normal Aging Skin. *Gerontology* 36, 28–35 (2004).
43. Derraik, J. G. B. et al. Effects of Age, Gender, BMI, and Anatomical Site on Skin Thickness in Children and Adults with Diabetes. *PLoS ONE* 9, e66657 (2014).
44. Sato, J., Yanai, M., Hirao, T. & Denda, M. Water content and thickness of the stratum corneum contribute to skin surface morphology. *Arch Dermatol Res* 292, 412–417 (2000).
45. Shuster, S., Black, M. M. & McVitie, E. The influence of age and sex on skin thickness, skin collagen and density. *Br J Dermatol* 93, 639–643 (1975).
46. Mattet, G., Magliaro, C., Pirone, A. & Alhuwalia, A. Decellularized Human Liver Is Too Heterogeneous for Designing a Generic Extracellular Matrix Mimic. *Artificial Organs* 3, e442 (2017).
47. Malda, J. et al. Of Mice, Men and Elephants: The Relation between Articular Cartilage Thickness and Body Mass. *PLoS ONE* 8, e57683 (2013).
48. Monteiro-Riviere, N. A., Bristol, D. G., Manning, T. O., Rogers, R. A. & Riviere, J. E. Interspecies and interregional analysis of the comparative histologic thickness and laser Doppler blood flow measurements at five cutaneous sites in nine species. *Investig Dermatol* 95, 582–586 (1990).
49. Jee, T. & Konvopoupolos, K. Skin viscoelasticity studied in vitro by microprobe-based techniques. *J Biomech* 47, 553–559 (2014).
50. Shen, Z. L., Kahn, H., Ballarini, R. & Eppel, S. J. Viscoelastic Properties of Isolated Collagen Fibris. *Biophys J* 100, 3008–3015 (2011).
51. Meliga, S. C. et al. The hyperelastic and failure behaviors of skin in relation to the dynamic application of microscopic penetrators in a microneedle model. *Acta Biomaterialia* 48, 341–356 (2017).
52. Smith, E. Histology of the Skin of the Elephant. *J Anat Physiol* 24, 493–503 (1890).
53. Synder, M. A. The Postnatal Development of Wool Follicles, Shedding, and Skin Thickness in Inbred Merino and Southdown-Merino Crossbred Sheep. *Aust. J. Biol. Sci.* 14, 141–156 (1961).
54. Nanan, M. & Mahmud, J. The effect of skin orientation on biomechanical properties. *J Mech Eng* (2015).
55. Chow, M.-Y. & Zhang, Y. Changes in the Mechanical and Biochemical Properties of Aortic Tissue due to Cold Storage. *JSE* 171, 434–442 (2011).
56. Banga, A. K. Transdermal and Intradermal Delivery of Therapeutic Agents. (CRC Press, 2011).
57. Prevost, T. P. et al. Dynamic mechanical response of brain tissue in indentation in vivo, in situ and in vitro. *Acta Biomaterialia* 7, 4090–4101 (2011).
58. Gepfken, A. & Margules, S. S. Are in vivo and in situ brain tissues mechanically similar? *J Biomech* 37, 1339–1352 (2004).
59. Walpole, S. C. et al. The weight of nations: an estimation of adult human biomass. *BMC Public Health* 12, 93 (2012).
60. Auriol, F. et al. Effects of short-time hydration on skin extensibility. *Acta Derm-Venereol* 73, 344–347 (1993).
61. Hewitson, T. D. & Darby, I. A. Histology protocols. (2010).
62. Meigel, W. N., Gay, S. & Weber, L. Dermal architecture and collagen type distribution. *Arch Dermatol Res* 259, 1–10 (1977).
63. Hayes, W. C., Keer, L. M., Herrmann, G. & Mockros, L. E. A mathematical analysis for indentation tests of articular cartilage. *J Biomech* 5, 541–551 (1972).
64. Takema, Y., Fujimura, T., Ohsu, H. & Imokawa, G. Unusual wrinkle formation after temporary skin fixation following by UVB irradiation in hairless mouse skin. *Experimental Dermatology* 5, 145–149 (1996).
65. Tchvialeva, L. et al. Skin Roughness Assessment. 341–348 (InTech, 2010). at http://www.intechopen.com/books/new-developments-in-biomedicalengineering/skin-roughness-assessment.
66. Fischer-Cripps, A. C. Introduction to Contact Mechanics. (Springer US, 2007). https://doi.org/10.1007/978-0-387-68188-7.
67. Delaine-Smith, R. M., Burney, S., Balkwill, F. R. & Knight, M. M. Experimental validation of a flat punch indentation methodology calibrated against unconfined compression tests for determination of soft tissue biomechanics. Journal of the Mechanical Behavior of Biomedical Materials 60, 401–415 (2016).
68. Prierat, B., MacManus, D. B., Murphy, J. G. & Gilchrist, M. D. Indentation of heterogeneous soft tissue: Local constitutive parameter mapping using an inverse method and an automated rig. Journal of the Mechanical Behavior of Biomedical Materials (2017). https://doi.org/10.1016/j.jmbbm.2017.03.033.
69. Lin, D. C., Shreiber, D. I., Dimitriadis, E. K. & Horkay, F. Spherical indentation of soft matter beyond the Hertzian regime: numerical and experimental validation of hyperelastic models. Biomach Model Mechanobiol 8, 345–358 (2008).
70. Wu, J. Z., Dong, R. G., Smutz, W. P. & Schopper, A. W. Nonlinear and viscoelastic characteristics of skin under compression: experiment and analysis. Biomed Mater Eng 13, 373–385 (2003).
71. Lin, D. C., Dimitriadis, E. K. & Horkay, F. Elasticity Models for the Spherical Indentation of Gels and Soft Biological Tissues. MRS Proc. 1060, 1060–LL05-07 (2011).
72. Beards, C. F. Engineering Vibration Analysis with Application to Control Systems. (Elsevier, 1995). https://doi.org/10.1016/B978-0-340-63183-4.X2900-5.
73. Ma, T., Har, M., Sougrat, R., Verbakat, J. M. & Verkman, A. S. Impaired Stratum Corneum Water Channel Hydration in Mice Lacking Epidermal Water Channel Aquaporin-3. J Biol Chem 277, 17147–17153 (2002).
74. Oznurlu, Y., Celik, I., Sur, E. & Telatar, T. Comparative Skin Histology of the White New Zealand and Angora Rabbits. Journal of animal and … 8, 1694–1701 (2009).
75. Jacobi, U. et al. Porcine ear skin: an in vitro model for human skin. Skin Res Technol 13, 19–24 (2007).
76. Oznurlu, Y., Celik, I., Sur, E. & Telatar, T. Comparative Skin Histology of the White New Zealand and Angora Rabbits. Journal of animal and … 8, 1694–1701 (2009).

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Author Contributions
J.C.J.W. performed the experiments. J.C.J.W., M.A.F.K. and M.L.C. prepared the figures. G.A.E., D.J.M., H.H., M.A.F.K. and M.L.C. provided scientific oversight on the experimental design, data analysis and interpretation. All authors contributed to the writing of the manuscript.

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