The Facial Adipose Tissue: A Revision

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Abstract

Recent advantages in the anatomical understanding of the face have turned the focus toward the subcutaneous and deep facial fat compartments. During facial aging, these fat-filled compartments undergo substantial changes along with other structures in the face. Soft tissue filler and fat grafting are valid methods to fight the signs of facial aging, but little is known about their precise effect on the facial fat. This narrative review summarizes the current knowledge about the facial fat compartments in terms of anatomical location, histologic appearance, immune-histochemical characteristics, cellular interactions, and therapeutic options. Three different types of facial adipose tissue can be identified, which are located either superficially (dermal white adipose tissue) or deep (subcutaneous white adipose tissue): fibrous (perioral locations), structural (major parts of the midface), and deposit (buccal fat pad and deep temporal fat pad). These various fat types differ in the size of the adipocytes and the collagenous composition of their extracellular matrix and thus in their mechanical properties. Minimal invasive (e.g., soft tissue fillers or fat grafting) and surgical interventions aiming to restore the youthful face have to account for the different fat properties in various facial areas. However, little is known about the macro- and microscopic characteristics of the facial fat tissue in different compartments and future studies are needed to reveal new insights to better understand the process of aging and how to fight its signs best.

Keywords
► facial fat
► fat compartments
► dermal adipose tissue
► subcutaneous adipose tissue

Comparing the histologic characteristics of the hypodermis amongst mammal species, different layered arrangements can be identified. Apes and dedicated furred mammal species display in their hypodermal fat layer a fibrous and/or muscular layer located between the dermis and the deep fascia (► Fig. 1). This specific layer is called the panniculus carnosus and separates the subcutaneous fat into a superficial and a deep fat layer (► Fig. 1A). In other mammals such as pigs (Sus scrofa domestica) that lack fur, this layer is absent and the total hypodermal fat layer is called panniculus adiposus (► Fig. 1B). In the latter arrangement, the perforating arteries can reach the skin and can regulate the dermal blood flow and thus skin temperature and nutrition. This different layered arrangement of the subcutaneous fat is believed to provide an evolutionary advantage to fur-less mammals (e.g., the human) as they are capable of adapting to different climate conditions based on
their advanced thermoregulatory capacity whereas other mammals cannot.\(^1\)

It is widely believed that the human hypodermal fat layer is absent of the *panniculus carnosus*. Investigating the three-dimensional (3D) functional anatomy of the human adipose tissue (AT)\(^2,3\) revealed a fascial plane composed of fibrous connective tissue and muscular fibers, which was identified in the hypodermal plane and can be regarded as the homologue to the *panniculus carnosus* in rodents.\(^2\)

In the head this separating layer is called *galea aponeurotica*; in the face, superficial musculo-aponeurotic system (SMAS); in the neck, platysma; in the abdominal region, *Scarpas’ fascia*; in the perineum, *Colles’ fascia*; and in the scrotum, *Dartos tunic*.

It has been shown that this hypodermal fascial arrangement subdivides the subcutaneous fat in the abdominal\(^4\) and facial\(^4\) regions into superficial and deep fat with different morphologic characteristics. To better understand this arrangement and the distribution of the white adipose tissue (WAT) in the face, we critically discuss here the recent insights in this field.

**Anatomy of Various Facial Fat Compartments**

**General Arrangement**

The face has a layered structure that can be described from superficial to deep as follows: *layer 1*: skin; *layer 2*: subcutaneous fat; *layer 3*: superficial musculoaponeurotic system (SMAS); *layer 4*: deep fat; and *layer 5*: deep fascia or periosteum. These five layers can be found throughout the face and are continuous from the head into the neck.\(^5\) From here two different compartmentalized layers of fat are continuously present in the face: superficial and deep. Of those, the superficial layer can be subdivided further into two different arrangements with different microstructure: in perioral, nasal, and in parts of periorbital areas versus the midface area. The deep layer itself can be differentially classified into the deep fat compartments and the buccal fat pad. These different facial fat arrangements have different functions and are characterized by different types of AT including its proper extracellular matrix (ECM).

**Superficial Fat Compartments (Layer 2)**

The subcutaneous fat layer contains dermal white adipose tissue (dWAT) and varies in thickness between individuals of different age and ethnicity. It is subdivided by *fibrous septae* into distinct compartments, which have been previously identified both in cadaveric\(^6-8\) and imaging studies.\(^9\) Within the walls of the bounding septae, nerves and vessels traversing this plane can be identified.\(^10\) Based on cadaveric dissections\(^5\) and on previous histologic analyses,\(^11\) two different types can be identified according to Ghassemi et al.\(^11\) Type 1 can be found in the medial and lateral midface and in parts of the periorbital region, as well as in the temple, forehead, and neck. Here, the adherence of the underlying structures to the skin is loose, and an easy separation of the skin from the underlying fat layer can be performed during anatomical dissections (►Fig. 2A). The WAT in these areas can be classified as structural WAT with a meshwork of fibrous septa enveloping lobules of fat cells that act as small pads with specific viscoelastic properties.\(^12\) Type 2 can be found in perioral and nasal areas, as well as in the area of the eyebrows. Here, a strong linkage is present between the facial muscles, the collagenous meshwork surrounding the adipocytes and the skin. A separation using blunt dissections is
difficult in these areas, whereas collagenous and muscular fibers directly insert into the skin and connect the skin to the underlying muscles of facial expression (Fig. 2B). The WAT here can be classified as fibrous with a meshwork of intermingled collagen and elastic fibers as well as muscle fibers. The latter reach up to the dermis of the skin and fat cells are interposed between this collagen fiber–muscle meshwork. The most prominent demarcation lines between these two subcutaneous arrangements include the nasolabial sulcus, the labiomandibular sulcus, and the submental sulcus (Fig. 3). Within the midface, the type 1 subcutaneous arrangement results in the formation of the nasolabial, medial, middle, lateral, and buccal subcutaneous fat compartments that have been previously introduced. It is worth mentioning that the medial subcutaneous fat compartment in layer 2 of the midface is also called the malar fat pad by some authors. On the forehead, three superficial fat compartments have been described previously and their description is in line with the arrangement of the fat layers in the midface.

**Deep Fat Compartments (Layer 4)**

The deep fat layer contains subcutaneous white adipose tissue (sWAT) and is separated from the more superficially located dWAT (layer 2) by the SMAS in the midface, platysma in the neck, and superficial temporal fascia in the temple. The orbicularis oculi and the frontalis muscle can also be identified within this layer as they are continuous with each other on the lower and the lateral forehead. The deep fat layer is likewise separated by fibrous septae into distinct compartments, and similar to the dWAT, these septae serve as transition pathways for the branches of the facial nerve or for branches of the facial artery and vein. Boundaries of deep fat compartments can also be formed by the origins of muscles of facial expression and their respective bony origins. The deep fat compartments have been recently described both in cadaveric and in imaging studies. The deep medial cheek fat compartment is enclosed by the levator labii superioris alaeque nasi and the levator anguli oris muscle and lies in close proximity to the infraorbital foramen. The triangular-shaped deep lateral cheek fat compartment is bounded medially by the facial vein, laterally by the zygomaticus major muscle, and superiorly by the zygomatic ligament. Medial to the deep medial cheek fat and lateral to the pyriform aperture the deep pyriform space can be identified, which previously was named Ristow-space. Another fat compartment (so-called deep nasolabial compartment) described both clinically and anatomically lies superficially to the levator labii superioris alaeque muscle within the premaxillary space but deep to the orbicularis oculi muscle. Superior to the zygomatic ligament but inferior to the orbicularis retaining ligament and lateral to the facial vein, the suborbicularis oculi fat (SOOF) compartment can be identified, which lies deep to the orbicularis oculi muscle. In the supraorbital region on the lateral margin of the orbit, the retro-orbicularis fat compartment (ROOF) is located, which is bounded medially by the neurovascular bundle emerging from the supraorbital foramen. In the lateral midface superficial to the masseter muscle, three distinct compartments can be identified, which provide avascular dissection pathways.
during face-lifting procedures: inferior, medial, and superior premassester compartments.\textsuperscript{15}

**Deepest Fat Compartments (Deep to Layer 5)**
Comparing with layers 2 and 4, the deepest fat layer is not continuous, but its fat compartments are few in numbers and the physiologic role of the fat within these compartments has not yet been fully elucidated. The buccal fat pad was first described in 1801\textsuperscript{24} and can be classified as such and is regarded as WAT of the deposit or metabolic type.\textsuperscript{12} It is located between the fascia of the buccinator muscle and the fascia of the masseter muscle in the buccotemporal space.\textsuperscript{25} The buccal fat is unique in humans and absent in most of ape species and has four extensions: buccal, pterygoid, pterygo-palatine, and temporal; the latter one is also known as the deep temporal fat pad of the temporal region and is found between the deep lamina of the deep temporal fascia and the temporalis muscle.\textsuperscript{26} However, another fat compartment in the deepest layer, which is much less well understood, is the superficial temporal fat pad (layer 6), and is found between the superficial and the deep lamina of the deep temporal fascia, superiorly to the zygomatic arch.\textsuperscript{26} While the superficial lamina of the deep temporal fascia is continuous anteriorly with the orbital septum,\textsuperscript{27} but not with the periosteum in the infraorbital region, it provides a space for fat within the prezygomatic space.\textsuperscript{9,23,27} This compartment lies deep to the SOOF and is separated from it by the superficial lamina of the deep temporal fascia.

**Facial White Adipose Tissue**

**Subcutaneous White Adipose Tissue**
As outlined previously, the facial sWAT can be further refined taking into account that the anatomically identified superficial and deep fat compartments are significantly different both morphologically and physiologically in respect to their WATs. Anatomically, the sWAT of the head and neck can be classified into two separate WAT depots. As it was mentioned previously, the interface between these WAT depots (superficial versus deep) in humans is less pronounced compared with rodents. Dynamically, deep sWAT is considered a slowly renewal tissue with characteristic turnover times of around 10 years.\textsuperscript{28,29}

**Dermal White Adipose Tissue**
Recently, it was demonstrated that adipocytes can behave dynamically in the superficial area of sWAT underlying the reticular dermis, where these cells can be involved in such processes as hair follicle cycling,\textsuperscript{30} wound healing,\textsuperscript{31} thermoregulation,\textsuperscript{32,33} and innate immune response of the skin.\textsuperscript{34} In all these processes, dermal adipocytes demonstrate the characteristic remodeling times of days to weeks. It was also assumed that these cells can be involved in spatially local cross-talk with skin cells leading to different inflammatory\textsuperscript{35–37} and pigmenitary\textsuperscript{35} skin conditions, as well as in skin aging.\textsuperscript{38} There are actually no genetic tools to selectively manipulate these subdepots since currently there is a lack of specific markers to differentiate between the superficial and deep adipocytes. At the same time, the ability of superficial adipocytes to undergo different phenotypic transformations\textsuperscript{35} (among them the recently discovered adipocyte-myofibroblast transition,\textsuperscript{39} which is now believed to play an important role in cutaneous fibrosis) clearly demonstrates the phenotypic plasticity of adipocytes from the dWAT layer. As a logical consequence, it was proposed to introduce dWAT as a special AT depot\textsuperscript{40,41} to differentiate between dermal adipocytes and adipocytes from a deep sWAT.

Furthermore, adipocytes from dWAT can locally react to internal and external factors with a quick modification in their number and cell volumes demonstrating either expansion or reduction of the dWAT layer (\textit{Fig. 4}). Resulting total modulation of WAT volume can many times exceed any effect that can be reached through a structural modification of the much thinner dermis. This evidently shifts the point of interest in noninvasive and minimal invasive aesthetic procedures from fibroblasts to adipocytes.\textsuperscript{42} At the same time, dWAT can be structurally and metabolically heterogeneous on the mesoscopic scale, which is an intermediate between the macroscopic scale of a single fat compartment and microscopic scale presented by single adipocytes or adipocyte clusters. Consequently, modification of this tissue by different internal or external factors can produce a spatial mosaic skin structure.\textsuperscript{36,37,43}

In humans, the dWAT does not produce the simple layers of adipocytes running almost parallel to the skin surface as in rodents but additionally demonstrate the protrusions into the dermis usually concentrated around the pilosebaceous units.\textsuperscript{35} A similar structure of dWAT was shown to be typical also for the porcine skin.\textsuperscript{44} Such dWAT structure provides spatially local clusters of dermal adipocytes, which, in an extreme case, can reach the papillary dermis and thus build the anatomical structures that can directly absorb the UV irradiation making dWAT equivalent to a potential target in the pathophysiology of the skin aging.\textsuperscript{38}

![Fig. 4](image_url) Imnunohistochemical staining with adipophilin of skin biopsy 19 days after the treatment with platelet-rich fibrin matrix showing differentiating dermal white adipocytes (red circles) within dermal white adipose tissue (dWAT). Note: Adipophilin is a protein associated with differentiating adipocytes.
Microscopic Heterogeneity of Adipose Tissue

Thus far, macroscopic anatomical description does not permit the differentiation of AT depots according to their structural components such as adipocytes and ECM. At the same time, recent experimental data have demonstrated that the AT structure is extremely heterogeneous on the microscopic and mesoscopic scales and that morphologic peculiarities and physiologic states of different AT depots are strongly dependent on the relative weighting of these components in a given AT volume. Adipocytes were for a long time considered to be the predominant structural component of every AT depot being responsible both for its physiologic properties and mechanical behavior as well as for its reaction to different external factors. This picture was strongly modified during the recent past as it became clear that other structural components of AT—especially, its ECM—are more important for the mechanical stability and renewal dynamics of ATs.

Adipocytes

Despite the fact that mature adipocytes occupy the main volume of AT, the number of these cells was estimated to be less than 20% of the total cellularity in ATs.5 Partial volume of every AT compartment depends on its cellularity and the characteristic adipocytes’ sizes, which vary significantly among different AT depots and have been shown to demonstrate sexual dimorphism.46–48 Adipocytes’ sizes in any AT depot have a typical bimodal distribution indicating the existence of at least two different subpopulations of small and large adipocytes.49 These distributions generally have high coefficients of variation of up to 50%. For example, it was shown that the patients with a body mass index (BMI) less than 25 kg/m² have 0.57 ± 0.23 μg of triglycerides per cell in femoral, 0.48 ± 0.15 μg in gluteal, and 0.41 ± 0.20 μg in abdominal areas, respectively. In patients with BMI greater than 30 kg/m², these values were shown to be 0.83 ± 0.18, 0.71 ± 0.23, and 0.78 ± 0.24 μg, respectively.46

Generally, the volume of any AT depot can be modulated in two different ways: through hyperplasia or hypertrophy, wherein hyperplasia is currently believed to be more characteristic in sWAT and hypertrophy is more often associated with the visceral fat depots.50 The hypertrophy of adipocytes was shown to be the first reaction to a high-fat diet,51 whereas the hyperplasia normally appears later and is connected with a prolonged excessive feeding. Physiologically small and large adipocytes are different,52 and it is believed that small adipocytes are mainly connected with development of new adipocytes from preadipocytes, which is required for good vascularization of AT depots. Adipocytes’ sizes can significantly differ in various body and facial areas,4 which can reflect different metabolic activities and fat structures in different facial AT compartments.

Extracellular Matrix in Adipose Tissues: Collagen

Adipocytes are imbedded in the extracellular matrix (ECM) of a special structure containing different types of collagen. The main function of this matrix was long believed to be the mechanical support: during AT expansion or involution, this matrix should be able to modify its structure to accommodate the altered adipocytes. This simple picture was substantially modified during the last years as it became clear that ECM in AT is not a passive scaffold but an active player in different morphologic and physiologic processes,53–56 and that its specific modification can even be a hallmark of metabolically changed adipocytes.

The role of collagen in ECM structure of ATs was long underestimated, until it was ascertained that different types of WAT have not only quantitative, but also dramatic qualitative differences in their collagen content. For example, AT of a hypertrophic type containing predominantly large adipocytes demonstrates also high content of collagens (Col) IV and VI, which are the nonfibrilляр collagens providing the pericellular fibrosis around the mature adipocytes.57,58 Content of Col VI in ATs demonstrates sexual dimorphism and varies significantly between different ethnic groups being higher in Asian Indians than in Caucasians.57 This fibrosis is supposed to restrict the unlimited expansion of adipocytes, which can lead to their damage and death, and this collagen was shown to be involved in different AT pathologies. Col VI content increases significantly with rising BMI; while the total fibrotic volume in sWAT is almost independent of BMI,59 the pericellular fibrosis increases with BMI by three to seven times 59,60. Contrarily to the pericellular fibrosis, the intercellular fibrosis exists in the form of thick collagen bundles consisting mainly of Col I and III that can penetrate the AT in different directions and can span many cell lengths.58 Both relative and absolute content of intercellular and pericellular fibrosis in different facial AT compartments can vary significantly.

Extracellular Matrix in Adipose Tissues: Hyaluronic Acid and Water Retention

Water content in an AT is normally less than 20%, whereas 14 to 16% of this water is located extracellular and only 2 to 6% intracellular. These values can be changed dramatically in a hypertrophic fat tissue, which normally demonstrates a significantly increased (up to 50% in extracellular space and up to four times in the pericellular areas) concentration of hyaluronic acid (HA).61

It was proposed that such high water retention around adipocytes prohibits the lipolysis, as the transport of water and glycerin in these cells should be realized through the same aquaporin 7 (AQP7) membrane channels.62 Though adipocytes have almost no cytoskeleton and thus appear nearly spherically in vitro, they can respond to a high osmotic pressure only with a limited expansion associated with a risk of the rupture of their cell membranes. This effect should force adipocytes to reduce the number of active AQP7 channels in the aqueous environment, which should consequently reduce the lipolytic activity of adipocytes. Such reduction in AQP7 channels in hypertrophic adipocytes was indeed observed in patients with high BMI values.63 Moreover, a depletion of HA content in AT through application of hyaluronidase was able to decrease the fat mass up to 35%, and this reduction was primarily connected with a reduction in the adipocytes’ sizes and not with a cell death, thus demonstrating a stimulation of the physiologic activity of adipocytes.64–66 HA found in a hypertrophic AT has...
a molecular weight of less than 15 kDa. It is known that the HA fragments of low molecular weight can stimulate the production of tissue fibrosis and that HA distribution in AT spatially correlates with appearance of fibrotic structures in this tissue. Moreover, HA was shown to interact with Col VI promoting its assembling in vitro. From this point of view, HA content can play a pivotal role in a local fibrosis and differential modification of mechanical properties of AT in different facial AT compartments.

**Microstructural Classification of Facial Adipose Tissue Depots**

Whereas the macroscopic compartmental classification of facial AT can be seen as state of the art in facial aging, a microstructural modification of different fat compartments in an aging face has not been described properly, although some authors indeed tried to make the differentiation.

The first attempt to describe the microscopic differences in the structure of various facial fat compartments taking into account both the morphology of adipocytes and the local collagen content was undertaken only recently. The background for this description was proposed in Sbarbati et al. who subdivided the sWAT depots using the results of transmission electron microscopy and scanning electron microscopy into three groups:

1. **Deposit (metabolic) WAT**: A nonlobular AT containing large adipocytes and characterized by extremely weak or even absent collagen network that does not wrap the single adipocytes.
2. **Structural WAT**: An AT containing large adipocytes, whereas each cell is evidently covered by a thin fibrous shell.
3. **Fibrous WAT**: An AT containing smaller adipocytes, whereas each cell is covered by a thick fibrous shell. This WAT was further subdivided into the lobular and nonlobular fibrous sub-types.

Based on these results, Bartossi et al proposed a classification of the fat compartments from the third medium of the face. These authors investigated the malar, periorbital, labial, nasal, and buccal fat pads using the same methods as Sbarbati and colleagues and have shown that

- Malar fat contains the lobules of large mature adipocytes homogeneously covered by thin collagen fibers as well as some newly formed adipocytes located near the capillaries (corresponding to the type 1 characterization of Ghassemi et al).
- Periorbital fat demonstrates the pronounced lobular structure with a dense network of thin fibers around the lobules producing the basket-like structures.
- Labial and nasal fat contains predominantly mature adipocytes embedded in a dense collagen matrix (corresponding to the type 2 characterization of Ghassemi et al).
- Buccal fat is characterized by large mature adipocytes that are not completely covered by a collagen network.

Following the classification proposed by Sbarbati et al, malar and periorbital fat pads were classified as WAT of the structural type, labial and nasal fat pads as WAT of the fibrous type, and buccal fat pad as WAT of the deposit type. This classification demonstrates that even the neighboring facial AT compartments can be structurally different, which can not only significantly influence the mechanical properties of the fat pads in single compartments but also mechanically stress the boundaries between these compartments produced by septa and thus cause a structural modification of the adjacent dermis area. However, it has to be noted that to date not all facial fat compartments have been investigated and detailed information is still missing. Future research will be needed to elaborate the precise characteristics of the remaining uninvestigated fat compartments like the superficial temporal fat pad or the fat within the prezygomatic space.

**Role of Adipocytes and Fibrotic Structures in Mechanical Properties of Adipose Tissue Depots**

Mechanical properties of AT depots play a pivotal role in the anatomy of the aging face. Taking into account that different facial AT compartments have different morphologic structures (especially concerning the content of pericellular and intercellular fibrosis and the average adipocytes’ sizes), it is important to understand how these parameters can modify the stiffness of specific AT depots.

Micromechanically, an AT depot can be described as a closed-cell foam structure. Because elastic shear modulus of triglycerides is much less than that of an AT, contribution of lipids into mechanical stiffness of sWAT can be neglected, even though they occupy the main tissue volume. Mechanical characteristics of such system are mainly dependent on the thickness of pericellular fibrotic structure containing Col IV and VI, which can vary significantly depending on the type of AT. Contrarily to the pericellular fibrosis, the intercellular fibrosis has a coarse-mesh structure with a typical unit size of several millimeters and its contribution to the total mechanical stiffness of AT depot is low. Mechanical strength of an AT depot without Col VI was recently measured to be approximately 50% lower than that of corresponding AT depot containing Col VI.

On the other hand, as it was shown by Kruglikov that local mechanical stiffness of AT depot is inversely dependent on the average cell size, being lower for the population of large adipocytes. Remarkably, this parameter is almost independent of the dispersion of adipocytes’ sizes. In other words, only the average cell size and not the cell size distribution is important for the mechanical stiffness of AT. Additionally, local appearance of small adipocytes (e.g., through induced adipogenesis or lipolysis) should improve the mechanical characteristics of the given AT depot. Moreover, an AT containing two different subpopulations of adipocytes of very different sizes can demonstrate a spatial heterogeneity of its mechanical properties.

According to the microstructural classification of the facial AT compartments provided in Bartossi et al., the “fibrous WAT” should have the highest content of pericellular fibrotic structures and thus the highest mechanical stiffness. On the
other hand, the “deposit WAT” is characterized by almost completely absent pericellular fibrotic structures and thus should have the worst mechanical properties.

Though every facial AT compartment undergoes a specific modification of its cellular and fibrotic components in aging, these changes can be very different even in neighboring facial compartments that can significantly influence the total facial aging appearance. For example, increase in the difference in mechanical modules in the neighboring compartments can produce a mechanical pressure on the septa between these facial areas leading to its deformation and consequent weakening in an aging face. Such deformation should be especially strong for the boundaries between AT compartments with sufficiently different pericellular structures. This effect can be clinically observed in the nasolabial and labiomandibular sulcus. Medial to both of the sulci, fibrous WAT can be identified, whereas lateral to them, structural WAT is present. During aging, the mechanical properties change and lead (along with other factors) to a bulging of the less mechanically stable tissue following gravity.

**Facial Aging on Mesoscopic Scale**

Present theories of skin aging almost completely exclude the involvement of AT in this process. At the same time, significant modification of dWAT was observed both in intrinsically and extrinsically aged murine skin. Reduction in the dWAT in a chronically UV-irradiated skin was shown to correlate with replacement of this volume by fibrotic structures indicating the involvement of recently revealed effect of adipocyte-myofibroblast transition, which mainly involves the dermal adipocytes. The appearance of skin wrinkles can be connected with a local disappearance of dermal adipocytes and deposition of fibrotic structures under the location of a skin depression.

Although sWAT and dermis appear anatomically as absolutely different tissues with a clearly defined boundary, there are several indications that these tissues strongly interact with each other. Co-culturing of adipocytes with fibroblasts demonstrated that hypertrophic (enlarged) adipocytes could significantly suppress the synthetic activity of fibroblasts through release of free fatty acids, whereas the small adipocytes had no such effect. Recently it was shown that the increment in sWAT correlates with decrease in elastic fibers’ content in the dermis and that this degradation can be connected with significantly elevated activity of matrix metalloproteinases. Consequently, not only the effects of chronological and photo-induced aging but also the expansion of sWAT can induce a relatively quick degradation of elastic fibers in the dermis, which, under normal conditions, should persist in this tissue for many decades. It can be speculated whether a local modulation of a superficial sWAT layer can lead to a corresponding response in an adjacent dermal area, leading to a change of its mechanical characteristics on the mesoscopic scale. Macroscopically, such negative adipocyte–fibroblast interaction can significantly influence the sagging severity in lower cheek and the formation of the nasolabial fold, both contributing to the formation of the aging face.

Another anatomical basis for wrinkles, which involves a specific fat depot, was recently proposed by Pessa et al. These authors found a spatial correlation between dedicated wrinkles (forehead crease, crow’s foot, and nasojugal crease) and the location of the big lymphatic vessels surrounded by the perilymphatic adipose tissue (PLAT). It is known that defective lymphatic transport can result in significant expansion of the AT; at the same time, it is not actually known whether modulation of PLAT can influence the local lymph transport. This raises the question whether the small, superficial PLAT depot located between well-known large facial AT compartments should be considered as a separate fat entity with the capability of influencing the process of facial aging.

**Soft Tissue Fillers Targeting Facial Adipose Tissue**

Application of soft tissue fillers (SFTs) is a broadly accepted minimally invasive antiaging procedure. Long-term skin improvement after injection of different types of SFTs was theoretically connected with activation of the neocollagenesis in dermis, which was criticized as an unrealistic mechanism. Independent of the filler type, SFTs are generally applied into the deeper located sWAT and thus it affects the AT structure. Moreover, it was shown that a midfacial STF injection demonstrates some beneficial effects also on the neighboring aesthetic units connected with another facial AT compartment.

Recently it was proposed to shift the main paradigm of SFTs from the neocollagenesis in the dermis to the spatial modification of the fat tissue and activation of adipose-derived stem cells (ADSCs). There are experimental results demonstrating that at least some SFTs (or their components) can stimulate the ADSCs in vivo. Whereas this item was not investigated with a scanning electron microscopy until now, it can be assumed that the sWAT might demonstrate the properties of the fibrous WAT after injection, thus improving the local mechanical properties in the injected area.

Better knowledge of facial fat compartments permits a more nuanced analysis of the aging face and better informs the next steps in restoring contour and facial balance, namely selecting optimal injection materials and the volumes required thereof. Injection materials have historically included fat as a filler and collagen-based products. Synthetic fillers are more contemporary alternatives, including HA-based fillers. Each STF is different in viscosity, stiffness, affinity for water, and in type and degree of cross-linking. Different STFs may be used, depending on their respective biophysical properties, to target specific AT compartments or subcompartments in varying depth and location. For example, STFs with higher stiffness and viscosity can be used to build volume and lift tissue in the midface or some deep locations in the face, where visibility is of less concern. STFs with less stiffness, which tend to diffuse or spread, are naturally more useful in superficial areas where the skin appearance is of main importance and the palpability is unwanted, such as the lower eyelids or lips.
Fat Grafts Targeting Facial Adipose Tissue

Injection of an autologous fat graft into the subdermal layer in facial regions was shown to provide significant modification of the dWAT layer, which appeared to have a richer microvascular bed after injection (Figs. 5–7). On the other hand, the pericellular fibrosis around adipocytes in injected area was significantly reduced, which can theoretically reduce the mechanical stiffness of the AT in this area. This can lead to a paradoxical situation: fat grafts can indeed immediately replace the lost volume in dWAT, but not really improve its mechanical properties.

The use of fat as a filler has been the subject of much study, including the investigation of the precise volume of injected fat required to achieve an optimal face “lift” effect. A study by Rohrich et al evaluated the “fill and lift” technique in 100 consecutive face lifts and noted that the following subcompartments are most important to fill prior to lifting the facial structures: the medial and middle superficial fat compartment, the deep medial and lateral cheek fat, and the deep nasolabial fat compartment. That study analyzed the synergistic effects of fat-grafting individual facial fat subcompartments in conjunction with either a “SMAS ectomy” or “SMAS stacking” technique, using the malar contour and nasolabial fold depth as key outcome measures. The average amount injected was 2 cc per fat compartment, with an average total of 12 cc per face (range: 8–14 cc), and an average of 2 cc in the nasolabial folds bilaterally. The study reported a malar projection increase of approximately 13.5% and malar tissue lift of approximately 12.2%. The study concluded that the “lift-and-fill” technique effectively addressed the core issue of facial aging, namely volume deflation, and that this technique required less tension to lift, resulting in a more natural appearance.

The perioral region having another WAT structure requires, however, a different approach during augmentation. This area is not typically undermined during a face lift, has superficial fat compartments, is the farthest point from the stretch of the lift, and is surrounded by adherent structures. During the aging process, there is a gradual loss of lip volume and support. Over time, lips become thinner, subcutaneous fat atrophies, nasolabial sulcus deepens, and the labiomandibular sulcus becomes more prominent. A study performed by Pezeshk et al evaluated and compared 65 patients who underwent rhytidectomy without perioral rejuvenation to 65 patients who underwent rhytidectomy with perioral rejuvenation using fat grafting. The pre- and postoperative photos were evaluated by three independent observers using a modified Fitzpatrick wrinkle scale (0: none, 1: mild, 2: moderate, 3: deep). Fat was harvested, centrifuged, and injected (1–2 cc into each perioral superficial compartment using a 22-gauge needle). Perioral injections.
were made in a fan-like manner to create a smooth lateral transition, with central expansion of the lips. Wrinkle scores decreased modestly (1.13–0.74 postoperatively) in the group with no perioral rejuvenation during rhytidectomy, while those patients with perioral rejuvenation improved more significantly (1.61–0.76 postoperatively). The authors emphasized that the face should be considered in three dimensions, taking into consideration the parameters of depth and width, and recommended that this area should be lifted and filled to address the volume depletion that occurs with aging.

Fat has unquestionable value as a filler in restoring volume, contour, and harmony to the face during rejuvenation surgery. Once selected as the appropriate filler to the task at hand, the focus shifts to proper harvesting and integration of the fat during autologous transfer. Multiple harvesting sites in the body have been described, such as the abdomen, flank, thigh, and knee. Recent studies evaluating adipocyte viability show that no harvest site is statistically better than another and that resorption rates after grafting are highly variable, between 20 and 90%.97

Subsequent studies have investigated inflammatory changes, as well as the changes that occur in the adipocytes at a cellular level that might influence viability. Suga et al98 investigated changes in mouse AT after ischemia/hypoxia caused by variably restricting blood supply to the AT. This model operated on the fact that the partial pressure in mice is similar to that of humans, measuring approximately 50 to 60 mm Hg. The study described a series of events in AT, beginning with induction of ischemia, followed by apoptosis of adipocytes, blood-derived cells, and endothelial cells on day 1 after harvesting. By days 3 and 7, the study described necrosis of the ATs, followed by infiltration of macrophages and phagocytosis. These ischemic changes were followed by regenerative actions, including angiogenesis, adipogenesis, and proliferation of cells. Accordingly, the authors suggest that supplementing a fat graft with ADSCs may reduce long-term graft atrophy and scar formation after severe ischemia occurring at the graft site as they were able to survive in the severe ischemia/hypoxia model.98

The use of ADSCs in conjunction with AT in fat grafting is a promising technique to provide volume and regenerate overlying, aged, and thinned skin. Further research on the role ADSCs in fat grafting may yield innovative and unprecedented application in treatment of the aging face.

**Conclusion**

The facial fat is a heterogeneous entity that can be subdivided into different anatomical compartments with each of them having different adipocyte morphology and different extracellular collagenous matrix. These different compositions provide unique and specific mechanical and histochemical properties to each of the respective compartments. Minimally invasive and surgical interventions aiming to restore the youthful face have to account for the different fat properties during therapeutic applications of STFs or fat grafting. However, still little is known about the macro- and microscopic characteristics of the facial fat tissue and future studies will be needed to reveal new insights to better understand the process of facial aging and how to fight its signs best.

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Statement

The facts and citations mentioned in the manuscript are reported to the best knowledge of the authors and potential deviations from the current understanding of the concepts presented should be considered in this narrative review.

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