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Dynamic mechanical properties of the tissue-engineered matrix associated with individual chondrocytes

BoBae Lee, Lin Han, Eliot H. Frank, Susan Chubinskaya, Christine Ortiz, and Alan J. Grodzinsky

Abstract

The success of cell-based tissue engineering approaches in restoring biological function will be facilitated by a comprehensive fundamental knowledge base of the temporal evolution of the structure and properties of the newly synthesized matrix. Here, we quantify the dynamic oscillatory mechanical behavior of the engineered matrix associated with individual chondrocytes cultured in vitro for up to 28 days in alginate scaffolds. The magnitude of the complex modulus (|E*|) and phase shift (δ) were measured in culture medium using Atomic Force Microscopy (AFM)-based nanoindentation in response to an imposed oscillatory deformation (amplitude ~ 5 nm) as a function of frequency (f = 1-316 Hz), probe tip geometry (2.5 μm radius sphere and 50 nm radius square pyramid), and in the absence and presence growth factors (GF, insulin growth factor-1, IGF-1, and osteogenic protein-1, OP-1). |E*| for all conditions increased nonlinearly with frequency dependence approximately f^1/2 and ranged between ~1-25 kPa. This result, along with theoretical calculations of the characteristic poroelastic relaxation frequency, f_p (~50-90 Hz) suggested that this time-dependent behavior was governed primarily by fluid flow-dependent poroelasticity, rather than flow-independent viscoelastic processes associated with the solid matrix. |E*(f)| increased, δ(f) decreased, and the hydraulic permeability, k, decreased with time in culture and with growth factor treatment.
This trend of a more elastic-like response was thought to be associated with increased macromolecular biosynthesis, density, and a more mature matrix structure/organization.

**Keywords**

Chondrocytes; AFM; Dynamic compression; Pericellular matrix; Growth factors; Nanomechanics; Tissue Engineering

**Introduction**

Cell-based tissue engineering holds great potential for therapies involving regeneration and/or replacement of damaged cartilage. Such approaches typically involve seeding cells (e.g., chondrocytes, stem cells) within macromolecular scaffolds or in scaffold-free environments and subjecting them to stimulatory biochemical and/or mechanical factors in vitro to promote synthesis and growth of neo-tissue for subsequent implantation in vivo. The ultimate goal and challenge is to produce a material with structural, biochemical and biomechanical properties similar enough to healthy cartilage so that upon maturation in vivo it can restore physiological function. To achieve this objective, it is important to understand the temporal evolution of the properties of the newly synthesized matrix. At early stages, matrix is formed around individual cells within the scaffold and these cell-matrix composites are often isolated from each other. As culture time increases, growth of neighboring cell-associated matrices leads to the formation of a more continuous neo-tissue which undergoes further evolution in structure and properties in vitro or in vivo. Throughout this process, the cell-associated matrix is important in facilitating cell signaling and mechanotransduction (Millward-Sadler 2000; Millward-Sadler 2004). Knowledge of the properties of the in vitro-generated matrix provides an assessment of the quality and ultimate success of a given tissue engineering approach and has great potential to be utilized for optimization.

Recently, we reported atomic force microscope (AFM)-based indentation of the newly synthesized cell-associated matrices of individual chondrocytes to quantify their quasi-static mechanical properties (Ng 2007). This approach provides an advantage over macroscopic testing of the entire tissue-engineered construct, which is complicated by the response of the cells and scaffold, a significant effect at early times (Buschmann 1995). In addition, such local measurements enable the quantification of cell-to-cell heterogeneity and the tissue engineering process in general, while macroscopic experiments yield a homogenized response. For chondrocytes cultured in fetal bovine serum (FBS)-supplemented medium in alginate scaffolds, the stiffness of the newly-formed cell-associated matrix increased from ~0.1 kPa on day 7 to 0.17 kPa by day 28 (Ng 2007), a value much less than that of the pericellular matrix of mature intact cartilage chondrons (~60 kPa) (Alexopoulos 2003), as well as cartilage extracellular matrix (~1 MPa) (Kim 1995). The use of insulin-like growth factor-1 (IGF-1) and osteogenic protein-1 (OP-1) supplementation significantly increased matrix stiffness ~25-fold to 4.15 kPa on day 28 compared to FBS culture on day 28, consistent with observed increases in total GAG accumulation, and indicative of a more mature structure (Ng 2007).

Several previous studies have focused on measurement of the time-dependent mechanical behavior of chondrocytes having no PCM e.g., using transient nanomechanical testing via micropipette aspiration (Trickey 2004), AFM-based nanoindentation (Darling 2007), dynamic osmotic loading (Chao 2005), optical tweezers (Huang 2003), and creep cytoindentation (Shieh 2006). In contrast, the present study investigates the dynamic oscillatory mechanical behavior (Mahaffy 2000; Alcaraz 2003; Mahaffy 2004; Park 2005; Smith 2005) of the cell-associated matrices of individual chondrocytes cultured in alginate scaffolds up to 28 days. Such time-dependent behaviors are important because tissue-engineered constructs implanted in vivo are
expected to undergo cyclic and impact loading that includes frequency components as high as 1 kHz, just as native cartilage does (Shepherd 1997). Given the known hydrated and porous microstructure of the tissue-engineered cell-associated matrix (DiMicco 2007), rate-dependent poro-viscoelastic processes are expected (Kim 2008). The objectives of this study were (1) to establish the experimental and theoretical methodologies for quantifying the dynamic mechanical properties of newly developed tissue-engineered matrix associated with single cells, (2) to investigate the temporal evolution of these properties with increasing time in 3D alginate gel culture, and (3) to quantify the effect of specific growth factors (IGF-1 and OP-1) on the dynamic mechanical functionality of the newly developed matrix.

Materials and Methods

Cell source and culture
Chondrocytes were obtained from femoral condyles of 1-2 week old bovine calves and seeded into 2% alginate scaffolds, as described previously (Ng 2007). Cell-seeded alginate beads were maintained either in high-glucose Dulbecco's Modified Eagle Medium (DMEM) with 10% FBS or in mini-ITS (29 ng/ml insulin, 2 μg/ml transferrin, 2 ng/ml sodium selenite) with the combination of IGF-1 (100 ng/mL) and OP-1 (100 ng/mL) (growth factor (GF) culture) (Loeser 2003). On days 7, 14, 21, and 28, cells were released from beads and suspended with high-glucose DMEM. Previous studies demonstrated that, under these same culture conditions, the cell-associated matrix was ~3 μm thick from day 7-28 (Ng 2007).

Dynamic oscillatory compression
For each group of 4-5 cells, an AFM probe tip was used to position each individual chondrocyte and its cell-associated matrix into one of the inverted pyramidal wells of a silicon substrate in DMEM (Fig. 1a). We were able to directly observe and choose each single cell on the substrate through the monitor via the optical microscope of AFM, as previously shown in Figure 1(c) of Ng et al (Ng 2007). A multimode Nanoscope IV AFM with PicoForce piezo (Veeco, Santa Barbara, CA) was then used to perform compression measurements on each cell-matrix composite within its well. Both pyramidal (Si₃N₄, cantilever spring constant, k~0.06 N/m, end radius, Rₜₐₚ ~50 nm, NP-20, Veeco) and then spherical colloidal probe tips (Silica beads from Bang Labs, #SS06N, tipless cantilever from Veeco, NP-020, k~0.06 N/m, Rₜₐₚ ~2.5 μm) were used sequentially to probe each cell of the given group of cells on the substrate. For each cell, the probe tip was centered manually (under visual control) over the well where the cell-matrix composite was placed. Script mode (Nanoscope IV software, V613r1) was programmed to apply a 1 μm ramp indentation at a displacement rate of 1 μm/s using both pyramidal and spherical colloidal probe tips (Ng 2007). A closed-loop scanner was used to apply the 1 μm ramp indentation and subsequently to maintain the indentation depth constant at 1 μm. Force relaxation at constant displacement was then measured during the subsequent 180-second interval (Fig. 1b,c). Only z-direction movement at zero x-y offset was allowed during the force mode AFM measurement to avoid crosstalk between AFM x-y and z-direction piezo displacements or x-y direction drift. After the force reached quasi-equilibrium within ~100 seconds, a sinusoidal displacement (5 nm amplitude) was superimposed on the 1 μm static offset displacement under closed loop control using an external wave generator (Rockland 5100) connected to the Nanoscope IV controller. The 5 nm displacement amplitude was determined from the measured z-piezo displacement. The magnitude and phase angle of the resulting force were measured at discrete frequencies of 1, 3.16, 10, 31.6, 100, and 316 Hz (Fig. 1c). The force magnitude was in the 0.01-0.1 nN range; at lower frequencies (<1 Hz) was below the AFM detection limit and software limitations prevented testing above 316 Hz. Measurements were repeated 3 times for each of the two probe tip sizes at the same single location of each cell-matrix composite. 4~5 cell-matrix composites from day 7, 14, 21, and 28

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cultures were tested with both probe tips. In addition, 3 day-0 cells were tested with nanosized probe tips (see Figure S1c).

Data analysis and statistical tests

Dynamic force and displacement data for both probe tips were analyzed using Fast Fourier Transforms to obtain the magnitude of the complex modulus, $|E^*(f)|$, and phase angle, $\delta(f)$, between measured force and applied displacement. Application of sinusoidal displacements even as low as 5 nm in amplitude, with either micron-sized or nano-sized probe tips, resulted in measured sinusoidal force waveforms having total harmonic distortion less than 5% over the entire frequency range of interest. $\delta(f)$ was corrected for the effect of hydrodynamic drag (Alcaraz 2002); the control experiment was performed on the Si substrate covered by culture medium to evaluate the phase angle of the control experiment using both probe tips (See Appendix D). For the micron-sized spherical probe tip, $|E^*(f)|$ was obtained from a Taylor series expansion of the Hertzian contact model (Johnson 1997; Mahaffy 2004):

$$|E^*_{\text{spherical}}(f)| = \frac{1 - v^2}{2 \sqrt{R d_0}} \frac{F(f)}{d(f)}$$

(1)

where $d_0$ is indentation depth at zero frequency, Poisson's ratio, $v$, was initially assumed to be 0.4 (Freeman 1994), $R$ is the effective radius of the probe tip and cell-matrix composite (i.e., $1/R = [1/R_{\text{spherical tip}}] + [1/R_{\text{cell-matrix composite}}]$). $F(f)$ is the magnitude of the force at frequency $f$, and $d(f)$ is the magnitude of the deformation at frequency $f$ during dynamic oscillatory compression (Ng 2007) (Fig. 1c). The Hertzian model approximation used in this study is sufficient for assessing relative trends in the temporal evolution of matrix stiffness with culture time, which is the primary focus of this study. The real $[E'(f)]$ and imaginary $[E''(f)]$ parts of the complex modulus (i.e., storage and loss moduli) are also reported, as well as $|E_{\text{pyramidal}}^*(f)|$ obtained using the pyramidal probe tip (see Appendix A).

To test our working hypothesis that culture duration and GF treatment significantly influence the dynamic properties of cell-matrix composites, each effect (culture duration, GF treatment, frequency and tip geometry) on $|E^*(f)|$ and $\delta(f)$ was tested using one-way ANOVA followed by Tukey’s post hoc test for comparisons. All data are expressed as mean±SEM, with $p \leq 0.05$ as statistically significant. Statistical analyses were performed using Systat-12.0 (Richmond CA).

Results

Effect of frequency

$|E^*|$ and $\delta$ increased nonlinearly with applied displacement frequency in both FBS (Fig. 2) and GF (Fig. 3) cultures at each time point. The effect of frequency on $|E^*|$ and $\delta$ was significant for both FBS and GF culture medium with both the spherical micron-sized tip (Figs. 2a,b, 3a,b) and the pyramidal nanosized tip (Figs. 4a,b, 5a,b, 6b) by one-way ANOVA (p<0.05). In FBS, $|E^*|$ and $\delta$ at 316 Hz were significantly different from that at all lower frequencies during the entire culture period (Fig. 2a,b, p<0.05). Similarly, for the nanosized tip, $|E^*|$ and $\delta$ at 316 Hz were significantly different from all lower frequencies during the entire culture period for both FBS and GF culture (Fig. 4a,b, 5a,b, p<0.05).

Effect of culture duration

$|E^*(f)|$, $E'(f)$, $E''(f)$ increased and $\delta(f)$ decreased with culture duration. The effect of culture duration on $|E^*(f)|$ and $\delta(f)$ was significant for both FBS and GF medium with the micron-sized (Fig. 2a,b, 3a,b) and nanosized tip (Figs. 4a,b, 5a,b) (one-way ANOVA, p<0.05). For GF

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culture, pair-wise comparisons showed that $|E^*(f)|$ obtained with the micron-size tip on day 14-28 at higher frequencies were significantly higher from that on day (p<0.05 for 316 Hz, p<0.005 for 100 Hz (not shown)). Fig. 6a shows the trend at 316 Hz between day 7 and day 21 in GF culture with the nanosized tip (p<0.05).

**Effect of GF treatment**

$|E^*(f)|$ and $E'(f)$ increased significantly with the addition of GF compared to FBS-supplementation using the micron-sized (Fig. 7a) and nanosized tip (Fig. 6c) (ANOVA, p<0.05). In GF culture, $\delta^*(f)$ was significantly lower than that in FBS at both low and high frequencies (1, 100-316 Hz) with the micron-sized (Fig. 7b) and nanosized tip (Fig. 6d) (p<0.05). Interestingly, the quasi-static force-displacement (indentation) curves of individual chondrocyte-matrix composites (Fig. S1a,b) demonstrated that GF-cultured samples exhibited a much higher indentation stiffness than FBS-cultured samples when utilizing a 1 $\mu$m indentation displacement. In contrast, while the dynamic stiffness $E^*(f)$ measured at 5 nm displacement amplitude increased significantly with GF-treatment, the increase was modest (~1.5-fold).

**The effect of tip geometry**

$|E^*(f)|$ was not significantly affected by tip geometry in this 28 day culture study. A possible interpretation is that after initial application of the ~1 $\mu$m deformation, the small amplitude oscillation (5 nm) involves approximately similar contact areas for both probe tips sizes once embedded within the matrix. In contrast, the matrix stiffness detected using larger 1 $\mu$m indentation deformation was much larger for the micron-sized colloidal tip (Fig. S1).

**Discussion**

An AFM-based single cell mechanics approach revealed that the dynamic mechanical properties of the newly synthesized tissue-engineered matrix-chondrocyte composite are frequency-dependent and change with culture duration and GF treatment. Dynamic oscillatory nanomechanics methods (Mahaffy 2000) were combined with an experimental protocol incorporating an inverted pyramidal Si well (Ng 2007) (Fig. 1a) to hold each cell-matrix composite in place, facilitating a more physiological-like shape. First, the mechanical contributions from the cell and matrix are discussed for the FBS and GF-supplemented systems. Following, the dynamic mechanical data of cells cultured in FBS are discussed, for which the matrix was much softer than that of the cell itself and, hence, the dynamic mechanical properties of the cell-matrix composite are dominated by the properties of the matrix. Scaling approximations are then used to identify the rate-dependent processes and energy dissipation mechanisms that underlie the observed dynamic behavior. Lastly, the effects of culture duration on indentation stiffness versus dynamic stiffness and the effects of GF stimulation are discussed.

**Individual contributions from cell and matrix**

Indentation data analyzed from the initial force-displacement (ramp) portion (Fig. 1b) of the experiments (Fig. S1a,c) showed that day 7-28 cell-matrix composites cultured in FBS were much more compliant than day 0 cells with no surrounding matrix, consistent with our previous study (Ng 2007). Finite element analysis of this system by Ng, et al. indicated that with FBS treatment, cell deformation was negligible compared to matrix deformation and, hence, the measured indentation stiffness was dominated by the matrix (Ng 2007). Hence, the dynamic mechanical properties reported here for the FBS culture conditions (Fig. 2) can be assumed to be dominated by the properties of the cell-associated matrix. However, for the GF cultures, nanoindentation data (Fig. S1b,d) show that the indentation stiffness of the cell-matrix composite increased substantially over 28 days, eventually approaching the stiffness of the
day-0 cell with no matrix. Therefore, under dynamic loading conditions, the dynamic mechanical properties of Fig. 3 likely include contributions from both the cell and matrix. We note that the Poisson’s ratio of the cell-matrix composite sensed by the probe tip may vary between day 7 and day 28, and may reflect the properties of both the cell and the cell-associated matrix (Alexopoulos 2005). We used the Poisson’s ratio of chondrocytes for our initial calculations (\( \nu = 0.4 \), (Freeman 1994). For conditions in which the matrix completely dominates the measured \(|E^*(f)|\), a value of \( \nu \) closer to that of the PCM (\( \nu = 0.04 \), (Alexopoulos 2005) may be more appropriate, which would increase the estimated \(|E^*(f)|\) by \( \sim 20\% \) in magnitude.

Mechanisms underlying dynamic frequency behavior

In general, the frequency dependence of \(|E^*|\), \(E'\), \(E''\), and \(\delta\) may be associated with fluid flow-dependent poroelasticity and/or flow-independent viscoelastic processes. To explore these possibilities, we first estimated the characteristic poroelastic relaxation time, \(\tau_p \sim [L^2/(Hk)]\) (also called the ‘gel diffusion time’) (Mow 1980; Grodzinsky 1981; Lee 1981; Soltz 2000; Morel 2004), where \(L\) is the characteristic length scale over which fluid flows, \(H\) is the elastic modulus, and \(k\) is the hydraulic permeability of the cell-associated matrix. \(L\) can be approximated as the radius of the probe tip, 2.5 \(\mu\)m (see Supplementary Appendix D for detailed discussion); \(H\) is approximated by quasi-static indentation modulus, \(~0.12\) kPa for day 14 and \(~0.17\) kPa for day 28 (Ng 2007); and \(k\) can be estimated from the measured GAG content (Ng 2007) to be \(~6.6 \times 10^{-12}\) m\(^4\)/N\(\cdot\)s for day 14 and \(~1.7 \times 10^{-12}\) m\(^4\)/N\(\cdot\)s for day 28 in FBS culture by using the unit cell model (Happel 1959; Eisenberg 1988) assuming that GAGs are the primary determinant of the permeability (see Appendix C). \(\tau_p\) of the newly developing matrix was thereby estimated to be \(~0.01-0.02s\) from days 14-28, corresponding to characteristic frequencies \(f_p \sim [1/\tau_p]\) of 50-90 Hz, which is within the 1-316Hz range of the testing frequencies of the present experiments and, in part, motivated the range of testing frequencies used.

Previous measurements and poroelastic models of the dynamic stiffness of native cartilage (Frank 1987; Kim 1995; Soltz 2000) have shown that \(|E^*|\) increases with \(f^{1/2}\) in the neighborhood of \(f_p\). \(|E^*|\) in the entire 1-316Hz range of Figs. 2a, 3a does, indeed, show an \(f^{1/2}\) dependence (\(R^2 > 0.94\), \(p < 0.001\)). This result, coupled with the value of \(f_p\) estimated above, suggests the importance of poroelastic behavior in the experimentally tested frequency range. Interestingly, for native cartilage explants, \(H\) is much larger and \(k\) much smaller than the cell-associated matrix, but the [Hk] product is on the same order (\(10^{-9}\) m\(^2\)/s). However, the much larger \(L\sim 1\) mm for cartilage explants tested in confined or unconfined compression results in frequencies \(f_p\) that are several orders of magnitude lower for the dynamic behavior of cartilage (0.001-0.01Hz) in confined or unconfined compression (Kim 1995; Buschmann 1999; Huang 2003). In contrast to this mm-length-scale testing, application of dynamic oscillatory compression to bovine cartilage explants using the micron-sized AFM probe tips again resulted in higher characteristic frequencies (\(f_p \sim 100\) Hz) (Han 2008), on the same order as that observed in the present study. For cartilage explants tested at the macroscale (Kim 1995), it was possible to perform experiments at frequencies \(f \gg f_p\) where it was observed that \(\delta(f)\) would begin to decrease with increasing frequency. It would therefore be very useful to obtain dynamic stiffness data for the cell-associated matrix at higher frequencies up to the 10 kHz range to more fully compare with theoretical predictions (Kim 1995). At present, however, AFM instrumentation limits data acquisition beyond a maximum frequency of \(~350\) Hz. The values of \(H\) and \(k\) estimated for our matrix may, indeed, reflect the structure and organization observed in previous studies (Lee 1998; DiMicco 2007) in which newly developing matrix revealed more diffuse and porous morphologies than that of freshly isolated, mature chondrons (Lee 1998). At the same time, intrinsic viscoelastic behavior associated with macromolecular constituents (Flory 1953) may also contribute to the frequency-dependent behavior of the newly developing matrix.
Effect of culture duration

The increase in $|E^*(f)|$ with time in culture is consistent with our previously measured increase in biosynthesis and deposition of GAG and collagen within the cell-associated matrix (Ng 2007). The decrease in $\delta(f)$ with culture duration corresponds to a decrease in energy dissipation during dynamic compression (i.e., a more elastic-like response), consistent with increased biomacromolecular deposition and altered matrix structure/organization (e.g., physical crosslinking, macromolecular entanglements). $k$ is also predicted to decrease with culture time (Appendix C), indicative of a denser matrix structure. It is hypothesized that the modest increase in $|E^*(f)|$ and the decrease in $\delta(f)$ with culture time compared to the quasi-static indentation stiffness may be due to the fact that the newly developing matrix is more sensitive to the higher compaction associated with the $1\mu$m indentation displacement compared to the much smaller ~5nm dynamic displacement amplitude.

Effect of GF treatment

The increase in $|E^*(f)|$ and decrease in $\delta(f)$ with GF treatment compared to FBS at the applied frequency range again suggests a more elastic-like response likely due to compositional and structural changes, including proteoglycan accumulation (Buschmann 1992; Mauck 2000; DiMicco 2007). The effect of GF treatment on $|E^*(f)|$ (Fig. 2a,3a) was modest compared to the more dramatic increase in the indentation stiffness (Fig. S1), suggesting that the effect of GF on functional mechanical properties is sensitive to the magnitude of the applied displacement.

In summary, our AFM-based approach has enabled the measurement of the poroelastic dynamic mechanical behavior of the newly developing matrix associated with individual chondrocytes, and can be applied generally to study the dynamic behavior of extremely compliant (~kPa) biological, porous, and hydrated systems at nm-length scales over a broad range of frequencies. The high resolution dynamic mechanical approach described here is able to discern fine differences in the development and maturation of the cell-associated matrix temporally and with the addition of growth factors. The methodologies reported here can thus be employed to assess maturation of matrix synthesized by primary chondrocytes and, additionally, by other cells such as stem cells undergoing chondrogenesis in applications for cartilage tissue engineering. It may also be possible to use this methodology to study the mechanobiological response of single chondrocytes (Shieh 2006) and stem cells to applied dynamic loads over a wide frequency range.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Abbreviations

- AFM: atomic force microscopy
- PCM: pericellular matrix
- ECM: extracellular matrix
- FBS: fetal bovine serum
- IGF-1: insulin-like growth factor
OP-1    osteogenic protein-1
GF       growth factor
IGF      standard error of mean

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Figure 1.
Dynamic oscillatory compression of individual chondrocytes and their associated matrix in FBS culture using a spherical tip: (a) Experimental setup: Chondrocyte-matrix composites in microfabricated silicon wells (radius of cell = R_{cell}, radius of probe tip = R_{tip}, matrix thickness = 3 μm, indentation depth = 1 μm, and small sinusoidal oscillation = 5nm; the detailed structure of the matrix has been described previously (Lee 1993; Dudhia 2005)), (b) Force (red) and indentation depth (blue) curve showing the initial indentation (depth, d_0 = 1 μm) and subsequent force relaxation exemplified by a day 14 chondrocyte, and (c) inset showing the applied dynamic sinusoidal deformation (d(f)) and resulting dynamic force (F(f)) shifted from the displacement by the phase angle δ(f).
Figure 2.
Dynamic mechanical properties of chondrocytes and their newly developing cell-associated matrix up to 28 days in FBS culture (10% FBS) using the spherical probe tip: (a) $E'(f)$, (b) $\delta(f)$, (c) $E''(f)$, and (d) $E''(f)$. n=number of cells, mean ± SEM; maximum SEM is ~4kPa for $E'(f)$ and ~6kPa for $E''(f)$, respectively.
Figure 3.
Dynamic mechanical properties of chondrocytes and their cell-associated matrix up to 28 days in GF culture (IGF-1+OP-1) using the spherical probe tip: (a) $E^*(f)$, (b) $\delta(f)$, (c) $E'(f)$ and (d) $E''(f)$. $n=$number of cells, mean ± SEM; maximum SEM is ~2kPa for $E'(f)$ and ~1kPa for $E''(f)$, respectively.
Figure 4.
Dynamic mechanical properties of chondrocytes and their newly developed matrix up to 28 days in FBS culture using the pyramidal probe tip: (a) $|E^*(f)|$, (b) $\delta(f)$, (c) $E'(f)$ and (d) $E''(f)$. n=number of cells, mean ± SEM; maximum SEM is ~2kPa for $E'(f)$ and ~3kPa for $E''(f)$, respectively.
Figure 5.
Dynamic mechanical properties of chondrocytes and their cell-associated matrix up to 28 days in GF culture (IGF-1+OP-1) using the pyramidal probe tip: (a) $|E'(f)|$, (b) $\beta(f)$, (c) $E'(f)$ and (d) $E''(f)$. n=number of cells, mean ± SEM; maximum SEM is ~2kPa for both $E'(f)$ and $E''(f)$. 
Figure 6.
Effect of culture duration and growth factor treatment on dynamic oscillatory properties using the pyramidal probe tip: (a) the effect of culture duration on $|E^*(f)|$ for GF culture chondrocytes (one-way ANOVA: culture duration), (b) the effect of frequency on $\delta(f)$ for GF culture chondrocytes (one-way ANOVA: frequency), (c) the effect of growth factor treatment on $|E^*(f)|$, and (d) the effect of growth factor treatment on $\delta(f)$ (one-way ANOVA: GF treatment).
Figure 7.
Effect of culture duration and growth factor treatment on dynamic oscillatory properties of chondrocytes and their newly developing matrix using the spherical probe tip: (a) the effect of growth factor treatment on $|E^*(f)|$ and (b) the effect of growth factor treatment on $\delta(f)$ (one-way ANOVA: GF treatment).