Human Meibum Lipid Conformation and Thermodynamic Changes with Meibomian-Gland Dysfunction

Douglas Borchman,1 Gary N. Foulks,1,2 Marta C. Yappert,3 James Bell,1 Emily Wells,1 Shantanu Neravetla,1 and Victoria Greenstone1

PURPOSE. Instability of the tear film with rapid tear break-up time is a common feature of aqueous-deficient and evaporative dry eye diseases, suggesting that there may be a shared structural abnormality of the tear film that is responsible for the instability. It may be that a change in the normal meibum lipid composition and conformation causes this abnormality. Principle component analyses of infrared spectra of human meibum indicate that human meibum collected from normal donors (Mn) is less ordered than meibum from donors with meibomian gland dysfunction (Md). In this study the conformation of Md was quantified to test this finding.

METHODS. Changes in lipid conformation with temperature were measured by infrared spectroscopy. There were two phases to our study. In phase 1, the phase transitions of human samples, Mn and Md, were measured. In phase 2, the phase transitions of model lipid standards composed of different waxes and cholesterol esters were measured.

RESULTS. The phase-transition temperature was significantly higher (4°C) for the Md compared with the Mn of age-matched donors with no history of dry-eye symptoms. Most (82%) of the phase-transition temperatures measured for Md were above the values for Mn. The small change in the transition temperature was amplified in the average lipid order (stiffness) at 33.4°C. The small change in the transition temperature was amplified in the average lipid order (stiffness) at 33.4°C. The average lipid order at 35.4°C for Md was significantly higher (30%, P = 0.004) than for Mn. The strength of lipid-lipid interactions was 72% higher for Md than for Mn. The ability of one lipid to influence the melting of adjacent lipids is termed cooperativity. There were no significant differences between Mn and Md in phase-transition cooperativity. However, there was a difference between Mn and Md in the minimum order or maximum order that Mn and Md achieved at very low and very high temperatures, respectively. The model wax studies showed that the phase transition of complex mixtures of natural lipids was set by the level of unsaturation. A double bond decreased the phase-transition temperature by approximately 40°C. The addition of a second CH=CH moiety decreased the phase-transition temperature by approximately 19°C. Unsaturated waxes were miscible with saturated waxes. When a saturated wax was mixed with an unsaturated one, the saturated wax disproportionately increased the phase transition of the mixture by approximately 30°C compared with the saturated wax alone. Cholesterol ester had little effect on the phase-transition temperature of the waxes. Model studies indicated that changes in the amount of lipid saturation, rather than the amount of cholesterol esters, could be a factor in the observed conformational changes.

CONCLUSIONS. Meibum lipid compositional changes with meibomian gland dysfunction reflect changes in hydrocarbon chain conformation and lipid-lipid interaction strength. Spectroscopic techniques are useful in studying the lipid–lipid interactions and conformation of lipid from individual patients. (ClinicalTrials.gov number, NCT00805452.) (Invest Ophthalmol Vis Sci. 2011;52:3805-3817) DOI:10.1167/iovs.10-6514

Meibomian gland and aqueous-deficient dry eye diseases affect 6 to 7 million people in the United States alone and can cause chronic and severe symptoms, as well as secondary changes that can lead to visual disturbance.1,2 Among many factors, maintenance of the integrity of tear lipids is essential to the prevention of dry-eye symptoms (see reviews by Foulks3 and others4–15). It is generally believed that the predominant cause of evaporative dry eye is meibomian gland dysfunction (MGD), although blink abnormalities also contribute to the disorder.16 The correlation between dry eye and an abnormal lipid layer of the tear film has been made by the use of interference microscopy17–22 and other techniques.23–25 Moreover, when the meibomian gland, which produces most of the tear film lipid, is dysfunctional, delivery of oil to the lid margin is reduced26–28 and dry-eye conditions occur.6,11 Changes in meibum composition that may contribute to and be diagnostic of dry eye symptoms have been reported.1,29–32

One of the earliest therapies for MGD has been to warm the eyelid to 45°C (113°F).33–35 and conformational changes observed in the hydrocarbon chains of Mn with temperature suggested that the observed increased delivery of meibum lipid with eyelid therapeutic heating could be related to the increased disorder in the packing of the hydrocarbon tails.36 Conformation is defined as the arrangement of the atoms in a molecule in space. The conformation of the lipid hydrocarbon chain changes with increasing temperature. When lipids are in an ordered state (at low temperatures) the hydrocarbon chains contain more trans rotomers (Fig. 1). Rotomers are conformations that can change with the rotation of a molecule. Order refers to a phase where lipids are in a more solid (but not completely) gel like phase. When lipids become disordered,

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they are in a fluid, liquid-crystalline phase and contain gauche rotomers. When lipids go from an ordered state to a disordered one, they are said to undergo a phase transition. The lipid phase transition is different from the melting of a solid, because the lipids are not completely solid and not completely liquid, but rather are more like a gel or liquid crystal. Phase transitions may be described by four parameters: the order of the most ordered and disordered state; the transition temperature at which half of the lipid molecules undergo a phase change; and the relative cooperativity. Cooperativity describes how a lipid molecule influences the order of neighboring lipids. The broader the phase transition, the smaller the absolute value for cooperativity. The phase-transition temperature is the temperature at which half the lipids undergo the phase transition. When the conformation of the hydrocarbon chains is all trans, the lipids are able to pack more closely together because the hydrocarbon chain is extended (Fig. 1) and Van der Waal’s forces are experimentally reproducible and were similar in multiple samples collected from the same person.39 Our studies showed that at ambient lid temperature, the lipid is approximately 37% ordered, between solid (gel phase) and liquid (liquid-crystalline phase).37 As the temperature increases from 25°C to 45°C, lipid delivery to the margins was observed to increase with a concomitant decrease in the refractive index,40 hydrocarbon disorder,36 and meibum lipid hydrocarbon motion.39 These findings suggest that hydrocarbon chain order and motion determine the delivery of meibum lipid from the meibomian glands to the lid margins and subsequently to the tear film.

Mn from the same person was approximately 40% less ordered at 35.5°C than was lipid extracted from tears, indicating that tears do not have the same lipid composition as Mn.36 In Mn and muscle, retina, and lens tissues, lipid saturation and lipid order at physiological temperature were linearly related to the lipid phase-transition temperature.43 Mn order and phase-transition temperatures decrease with age, a trend that may be attributable to lipid compositional changes.40,42 Our infrared spectral studies, along with principal component analysis, enable the quantification of the variance among the lipid spectra, which allowed us to identify and quantify protein associated with human meibum.43 The spectra were used to discriminate between Mn and meibum from donors with MGD (Md) with an accuracy of 93%.43 More important, this high degree of accuracy shows that the infrared spectra contain compositional and structural information about the changes that occur with meibomian gland dysfunction. Md was found to contain more protein and relatively less methyl groups (CH₃) and cis double bonds (cis=CH) than did Mn.37 The amount of protein was confirmed from relative infrared band intensities.37

We tested Md both before and after 1 month of therapy with topical azithromycin44 and determined that this therapy reduces elevated phase-transition temperature (a general biomarker) and lipid order (another biomarker) of abnormal secretions to normal levels. Md conformation was measured with temperature to characterize its thermodynamic properties for comparison with those of Mn.

MGD is defined as “a condition of progressive obstruction due to ducal hyperkeratinization or inspissation of secretion.”5 Until this study, inspissation (thickening) of meibum lipid had been assessed only qualitatively. One would expect that the lipid-lipid interactions would be greater in thicker secretions. In this study, the thickening of meibomian secretion was quantified by infrared spectroscopy. This technology has been applied to assessment of the molecular structure/conformation and packing of meibum lipid hydrocarbon chains.36–40,44

In phase 1 of our study, infrared spectroscopy was used to characterize the biophysical and thermodynamic properties of Md in terms of lipid order, lipid-lipid interaction strength and phase-transition temperature parameters. In phase 2 of our study, model wax and cholesterol ester mixtures were studied to determine the major compositional factors that contribute to changes in the thermodynamic properties of meibum.

Methods

Materials

Silver chloride windows for infrared spectroscopy were obtained from Crystran Ltd. (Poole, UK). Wax esters, oleyloleate (OO) and palmitoleate (PO), palmitoyloleate (PO), palmitolactate (PT), sterylpalmitate (SP), and the cholesterol ester cholesteryl palmitate were purchased from the Sigma-Aldrich (St. Louis, MO).

Diagnosis of Normal and MGD Status

Normal status was assigned when the patient’s meibomian gland orifices showed no evidence of keratinization or plugging with turbid or...
thickened secretions, and no dilated blood vessels were observed on the eyelid margin. Normal donors did not recall having dry eye symptoms. The diagnosis of MGD was made according to the criteria of Foulks and Bron.26 Plugging of the meibomian glands in at least 5 of 10 orifices in the central portion of the upper eyelid was a requirement for diagnosis. The character of meibomian gland–expressed secretion had to be turbid, turbid with clumps, or pastelike. Inflammation of the eyelid margin, as evidenced by swelling of the eyelid margin and 2+ vascular injection of the posterior lid margin, was necessary for diagnosis. The presence of telangiectasia of the posterior eyelid margin was confirmatory of chronic disease but was not required for study enrollment. Inclusion criteria were the presence of symptomatic meibomian gland dysfunction in subjects between 18 and 80 years of age who were not taking systemic or topical antibiotics and were not using topical anti-inflammatory medications. Exclusion criteria were a history of allergy to azithromycin, altered lid anatomy (with the exception of meibomian gland dysfunction), wearing eye makeup, inability to complete the prescribed therapy and follow-up. Dietary history was not assessed, but none of the subjects were taking systemic or topical antibiotics and were not using any further manipulation or extraction. For phase 2, the study of model waxes, standard wax was dissolved in tetrahydrofuran/methanol (3:1, vol/vol), and approximately 1 mg was applied to an AgCl infrared window. The solvent was evaporated under a stream of nitrogen gas, and the window was placed in a hypotilizer for 4 hours to remove all traces of solvent.

Fourier Transform Infrared Spectroscopy
For phase 1, the study of human meibum, expressed meibum was placed directly from the platinum spatula onto an AgCl window without any further manipulation or extraction. Infrared data analysis was then performed (GRAMS/386 software; Galactic Industries, Salem, NH). The frequency of the CH2 band near 2852 cm−1, because measurement of the symmetric stretch 1 band rather than the near 2954-cm−1 asymmetric CH2 stretching band is useful for estimating the content of trans and gauche rotamers in the hydrocarbon chains. Although the 2954-cm−1 asymmetric CH2 stretching band is useful for measuring phase-transition parameters, we chose to use the 2850-cm−1 band rather than the near 2954-cm−1, because measurement of the asymmetric band frequency is complicated by the adjacent CH3 symmetric stretching band near 2995 cm−1 and the CH2 symmetric stretching band near 2852 cm−1. The symmetric stretch (Θsym) was calculated by first baseline leveling the OH-CH stretching region be-

### Table 1. Grading of Clinical Symptom in Donors with MGD

<table>
<thead>
<tr>
<th>Sign</th>
<th>Mean Grade</th>
<th>Grade 0</th>
<th>Grade 1</th>
<th>Grade 2</th>
<th>Grade 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Itching</td>
<td>1.1 ± 0.2</td>
<td>None</td>
<td>Awareness</td>
<td>Desire to rub</td>
<td>Frequent rub</td>
</tr>
<tr>
<td>Fbs</td>
<td>1.4 ± 0.2</td>
<td>None</td>
<td>Awareness</td>
<td>Desire to rub</td>
<td>Desire to close lid</td>
</tr>
<tr>
<td>Dryness</td>
<td>0.7 ± 0.2</td>
<td>None</td>
<td>Awareness</td>
<td>Need for drop</td>
<td>Frequent drops</td>
</tr>
<tr>
<td>Burning</td>
<td>0.7 ± 0.2</td>
<td>None</td>
<td>Awareness</td>
<td>Need to rub</td>
<td>Frequent rub</td>
</tr>
<tr>
<td>Swelling</td>
<td>0.73 ± 0.08</td>
<td>None</td>
<td>Noticeable</td>
<td>Obvious</td>
<td>Decrease in aperture width</td>
</tr>
</tbody>
</table>

Values are average ± SEM. Fbs, foreign body sensation.

### Table 2. Grading of Clinical Signs in Donors with MGD

<table>
<thead>
<tr>
<th>Sign</th>
<th>Mean Grade</th>
<th>Grade 0</th>
<th>Grade 1</th>
<th>Grade 2</th>
<th>Grade 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plugged, n</td>
<td>9.0 ± 0.4†</td>
<td>None</td>
<td>Pink</td>
<td>Light red</td>
<td>Bright red</td>
</tr>
<tr>
<td>Lid margin redness</td>
<td>2.0 ± 0.2</td>
<td>None</td>
<td>Rounded</td>
<td>&gt;1 mm</td>
<td>&gt;2 mm</td>
</tr>
<tr>
<td>Swelling</td>
<td>0.9 ± 0.2</td>
<td>None</td>
<td>Round</td>
<td>Light pressure</td>
<td>Moderate pressure</td>
</tr>
<tr>
<td>Plugging</td>
<td>3.2 ± 0.2</td>
<td>Touch</td>
<td>Turbid</td>
<td>Turbid with clumps</td>
<td>Solid paste</td>
</tr>
<tr>
<td>Mg secretion</td>
<td>3.2 ± 0.2</td>
<td>Clear</td>
<td>Light pressure</td>
<td>Moderate pressure</td>
<td>Occluded</td>
</tr>
<tr>
<td>TBUT</td>
<td>6.0 ± 0.3†</td>
<td>None</td>
<td>Transparent</td>
<td>Transparent</td>
<td>Transparent</td>
</tr>
</tbody>
</table>

Values are average ± SEM.† TBUT, number of seconds to first breakup following complete blink.
between 3500 and 2700 cm⁻¹. The center of mass of the CH₂ symmetric stretching band was calculated by integrating the top 10% of the intensity of the band. The baseline for integrating the top 10% of the intensity of the band was parallel to the OH-CH region baseline.

Lipid CH₃ groups in the hydrocarbon chains are present as gauche rotomers, prevalent in disordered hydrocarbon chains, or trans rotomers, more abundant in ordered hydrocarbon chains (Fig. 1). Thus, lipid hydrocarbon chain order may be evaluated in terms of the relative amount of CH₃ trans rotomers. The frequency of the CH₂ symmetric stretch, ν̃_{sym}, is dependent on the amount of trans or gauche rotomers and has been used to characterize lipid conformational order (see Borchman et al.36,40). The data for percentage of trans rotomer were used to calculate the phase-transition enthalpy and entropy to convert a mole of gauche rotomers at 33.4°C and dividing by 100 to estimate the enthalpy of the transition and then converting ν̃_{sym} to the percentage of trans rotomers, a measure of lipid conformational order (see Borchman et al.36,40). Since rotomers are either in trans or gauche conformations, phase transitions can be described by a two-state sigmoidal equation, as described by Borchman et al.36 Lipid order at 33.4°C was calculated by extrapolating the ν̃_{sym} at 33.4°C from the fit of the phase transition and then converting ν̃_{sym} to the percentage of trans rotomers, a measure of lipid conformational order (see Borchman et al.36,40). The data for percentage of trans rotomer were used to calculate the phase-transition enthalpy and entropy from the slopes of Arrhenius plots, as described in Borchman et al.36,40. Arhenius plots from tear–lipid phase transitions were linear, with correlation coefficients greater than 0.998.

Because we calculated the percentage of trans rotomers at 33.4°C and calculated the enthalpy and entropy to convert a mole of gauche rotomers to trans rotomers, we could calculate the strength of lipid–lipid interactions at 33.4°C by multiplying lipid order (%) by the enthalpy of the transition and dividing by 100.

**Statistics**

Data are presented as the average ± the SEM. Significance was determined with the Student’s t-test or the correlation coefficient from the linear regression best fit. Differences reaching P < 0.01 were considered to be statistically significant.

### Table 3. Donor Data

<table>
<thead>
<tr>
<th></th>
<th>Normal</th>
<th>Meibomian Gland Dysfunction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>81/26</td>
<td>62/31</td>
</tr>
<tr>
<td>Caucasian</td>
<td>78/25</td>
<td>78/59</td>
</tr>
<tr>
<td>Black</td>
<td>3/1</td>
<td>12/6</td>
</tr>
<tr>
<td>Hispanic</td>
<td>0/0</td>
<td>2/1</td>
</tr>
<tr>
<td>Asian</td>
<td>16/5</td>
<td>0/0</td>
</tr>
<tr>
<td>Unknown</td>
<td>5/1</td>
<td>8/4</td>
</tr>
<tr>
<td>Total number of donors*</td>
<td>32</td>
<td>50</td>
</tr>
</tbody>
</table>

* Data are the percentage/number of the total group. A few donors donated as many as five times.

### Table 4. Lipid Phase-Transition Parameters

<table>
<thead>
<tr>
<th>Phase-Transition Parameter</th>
<th>Normal (Age-Matched)</th>
<th>MGD</th>
<th>Significance (P*)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minimum frequency, cm⁻¹</td>
<td>2849.90 ± 0.09</td>
<td>2849.68 ± 0.06</td>
<td>0.09</td>
</tr>
<tr>
<td>Maximum frequency, cm⁻¹</td>
<td>2853.6 ± 0.1</td>
<td>2853.48 ± 0.07</td>
<td>0.41</td>
</tr>
<tr>
<td>Cooperativity</td>
<td>-9.2 ± 0.5</td>
<td>-8.5 ± 0.4</td>
<td>0.40</td>
</tr>
<tr>
<td>Phase transition</td>
<td>28.9 ± 0.4</td>
<td>32.2 ± 0.5</td>
<td>0.002</td>
</tr>
<tr>
<td>Temperature, °C</td>
<td>9.2 ± 0.5</td>
<td>-8.5 ± 0.4</td>
<td>0.40</td>
</tr>
<tr>
<td>Enthalpy, kcal/mol</td>
<td>18.0 ± 15</td>
<td>15.7 ± 7</td>
<td>0.15</td>
</tr>
<tr>
<td>Entropy, kcal/mol/deg</td>
<td>0.56 ± 0.05</td>
<td>0.51 ± 0.02</td>
<td>0.29</td>
</tr>
<tr>
<td>Order % at 33.4°C</td>
<td>35 ± 2</td>
<td>47 ± 2*</td>
<td>0.004</td>
</tr>
<tr>
<td>Donors, n</td>
<td>13</td>
<td>49</td>
<td></td>
</tr>
</tbody>
</table>

* Order values are predicated on meibum from 50 donors.

**RESULTS**

### Phase 1: Human Meibum Study

**Human Material.** Human donors of meibum were recruited from the members and associates of DB’s laboratory and patients of GNF’s at the Kentucky Lions Eye Center and the Veterans Affairs Hospital, Louisville Kentucky. Donor data are presented in Table 3. The number of donors is different in Tables 3 and 4, because Table 3 lists all donors and Table 4 lists only age-matched normal donors.

### Lipid Phase Transitions and the Infrared CH Stretching Region.** The CH₂ stretching bands were predominant in the infrared spectra of lipids due to the large number of CH₂ groups in their hydrocarbon chains (Fig. 2). The CH stretching region is composed of six major bands.39 In this study, we used ν̃_{sym} near 2850 cm⁻¹ to estimate the trans to gauche rotomer content of the hydrocarbon chains (spectrum a). The spectrum of lysozyme (d) was scaled to the amide II band (a) to show that even if all the intensity of the band marked amide II in (a) were from protein, there would be no significant overlap with the lipid CH stretching bands.

![Figure 2](https://iovs.arvojournals.org/pdfaccess.ashx?url=/data/journals/iovs/933250/)
respectively; the transition temperature at which half of the lipid molecules underwent a phase change; and the relative cooperativity. The broader the phase transition, the smaller the absolute value of the cooperativity. Cooperativity describes how the order of a lipid influences that of neighboring lipids.

The four phase-transition parameters necessary for defining the phase transition and other parameters calculated from the defining parameters are listed in Table 4. The phase-transition temperature was significantly (4°C) higher ($P < 0.01$) for Md than for Mn of age-matched donors with no history of dry eye symptoms (Table 4).

The individual phase-transition temperatures versus age are shown in Figure 3A. Only 18% of the phase-transition temperatures measured for Md were within the 95% confidence limits for Mn (Fig. 3A). Note that the lowest values for the phase-transition temperature decreased with age at the same rate as the values for Mn reported previously (Fig. 3A).38 None of the nine phase-transition temperatures measured for Mn was above the average phase-transition temperature for Md. There were no significant differences in the other three phase-transition parameters between Mn and Md (Table 4).

Lipid order was measured close to but not exactly at the temperature of the eye lid, 33.4°C, so lipid order at exactly 33.4°C was calculated by extrapolating the $\tilde{\nu}_{\text{sym}}$ at 33.4°C from the fit of the phase transition and then converting $\tilde{\nu}_{\text{sym}}$ to the percentage of trans rotomers (Table 4).38 The average lipid order at 33.4°C for Md was significantly (30%; $P < 0.0001$) higher than for Mn (Fig. 3B, Table 4). The individual lipid orders at 33.4°C versus age are shown in Figure 3B. Note that the lowest Md values for the lipid order at 33.4°C decreased with age at the same rate as the values reported for Mn reported previously (Fig. 3B).40 No sex- or race-related differences between the phase-transition parameters of Md were evident.

The large interdonor distribution of parameters (Fig. 3) was not due to large variations in the precision of testing protocols. Phase-transition temperatures spanned 12°C from sample to sample, which is much larger than the ±2°C precision of the measurements (Fig. 3). The variability from sample to sample must have been due to differences in lipid composition differences. The enthalpy, $157 \pm 7$ kcal/mole, and entropy, $0.51 \pm 0.02$ kcal/mole/deg, of the phase transitions of Md were not significantly different from those of Mn (Table 4) and did not change with age. The strength of lipid–lipid interactions was 72% higher for Md ($43 \pm 6$ kcal/mole) than for Mn ($74 \pm 4$ kcal/mole).

The order versus phase-transition temperature data (Fig. 4) were fit by linear regression analysis, and the slope of the fit was 0.26, with an intercept of 19.8 ($r = 0.76, P < 0.005$). This result indicates that the order of human meibum at 36°C is directly related to the phase-transition temperature within the range of phase-transition temperatures measured. When the lipid reaches its maximum or minimum order, the phase-transition temperature is at a critical point, above or below which the lipid order will not change.

**Phase 2: Model Lipid/Wax Study**

Elongated waxes are tube shaped, forming multilamellar monolayers that are closely packed (2.5–3.2 Å; Fig. 1).54,55 When waxes are ordered (stiff, or unmelted), the hydrocarbon chains align in a trans conformation and pack in a perpendicular orthorhombic crystal structure (Fig. 1, top).54 Structural analysis has not been done on disordered (fluid, or melted) waxes that take on gauche conformations. In this study, we examined...
the influence of double bonds and cholesterol ester on wax mixtures, since wax is the major component of meibum.56

The Phase-Transition Temperature and the Infrared CH Stretching Region (between 3100 and 2700 cm\textsuperscript{-1}). Hydrocarbon order-to-disorder phase transitions were measured for four model standard wax esters over a temperature range of −20°C to 70°C (Fig. 5A). Hysteresis was noted in all samples. Compared to the heating curves, the phase-transition temperatures for the cooling curves were approximately 7°C lower for the unsaturated wax esters oleyloleate (OO) and palmityloleate (PO) (Table 5). Hysteresis in the phase transitions of the saturated waxes palmityloleytrate (PP) and sterylpalmitate (SP) was less (≈0.8°C) than that for the unsaturated waxes, but measurable (Fig. 5B, Table 5). Compared to PP, an additional \( \text{CH}_2 \)–\( \text{CH}_2 \) moiety in the palmitylohydrocarbon chain of SP raised the melting temperature (Tm) approximately 5°C (Table 5). A double bond in PO decreased the phase-transition temperature of SP by approximately 40°C. In comparison, the addition of a second CH\( = \)CH moiety in OO decreased the phase-transition temperature of PO by approximately 19°C.

The unsaturated waxes OO and PO were miscible with the saturated waxes PP and SP, as is evident by there being only one phase transition (Figs. 5A, 5C). When a saturated wax, PP, was mixed with an unsaturated one, PO, the saturated wax, disproportionately increased the phase transition of the mixture by approximately 30°C compared with PO alone (Table 5, Figs. 5A). In comparison, PO lowered the phase-transition temperature of the mixture by only approximately 5°C compared with PP alone. Similarly, when the saturated wax SP was mixed with an unsaturated one, OO, the saturated wax disproportionately increased the phase-transition temperature of the mixture by approximately 50°C, compared with OO alone (Table 5, Fig. 5C). In comparison, OO lowered the phase-transition temperature of the mixture by only approximately 6°C compared with SP alone. The cooperativity of the PO/PP or OO/SP mixtures was lower than that of PP or SP alone (Table 5).

Cholesterylpalmitate (20 mole percent) increased the Tm of the equimolar OO/SP wax mixture by slightly more than 1°C in the heating and cooling curves (Fig. 5D, Table 5). At 24°C, the

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Cholesterylpalmitate (20 mole percent) increased the Tm of the equimolar OO/SP wax mixture by slightly more than 1°C in the heating and cooling curves (Fig. 5D, Table 5). At 24°C, the

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**Figure 5.** (A) Wax phase transition curves for OO oleyloleate, PO palmityloleate, PP palmityloleate, SP sterylpalmitate. (●) Heating curve; (○) cooling curve. Lines are the fit of the data to equation 1 in Borchman et al.56 The lower the CH\( _2 \) symmetric stretching frequency, the more ordered the wax hydrocarbon chains. (B) Melting curve for sterylpalmitate. The scale is enlarged from (A) so that hysteresis between the cooling (○) and heating (●) curves can be observed. (C) Equal molar mixture of sterylpalmitate and oleyloleate (●) heating and (○) cooling curve. Sterylpalmitate: oleyloleate:cholesterylpalmitate 2:2:1 (m:m:m) where m is moles. (▲) heating curve, (△) cooling curve. (D) Enlargement of the scale in (C).
Replicate runs were combined, and then the phase-transition parameters were calculated.

Stearylpalmitate/oleyloleate/cholesterylpalmitate Palmitylpalmitate/palmityloleate, 1:1, m:m

that the hydrocarbons of cholesterol alone were ordered.

Branes (Fig. 6). The data were fit to the line with a slope of 

meibum, which is predominantly wax and other natural mem-

saturate, and similar to the linear relationship between satu-

meibum donor data are presented 

phase-transition temperature measured for human 

 satiation, and similar to the linear relationship between satu-

molecular membrane, SR S slow twitch rabbit muscle sarcoplasmic reticu-

TABLE 5. Phase-Transition Parameters

<table>
<thead>
<tr>
<th>Sample</th>
<th>Transition Temp. (°C)</th>
<th>Minimum Frequency (cm⁻¹)</th>
<th>Maximum Frequency (cm⁻¹)</th>
<th>Cooperativity (No Units)</th>
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<tr>
<td>Oleyloleate</td>
<td>Heating 0.5 ± 0.3</td>
<td>2849.84 ± 0.03</td>
<td>2853.11 ± 0.04</td>
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<td></td>
<td>Cooling −10 to −4</td>
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<td></td>
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<tr>
<td>Palmitoyloleate</td>
<td>Heating 19.96 ± 0.02</td>
<td>2849.1 ± 0.1</td>
<td>2853.7 ± 0.2</td>
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<td></td>
<td>Cooling 15.1 ± 0.1</td>
<td>2848.6 ± 0.6</td>
<td>2853.5 ± 0.4</td>
<td>45 ± 16</td>
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<tr>
<td>Palmitoylpealmitate</td>
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<td>2849.31 ± 0.05</td>
<td>2854.0 ± 0.1</td>
<td>480 ± 178</td>
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<td>Cooling 51.06 ± 0.01</td>
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<td>2853.91 ± 0.04</td>
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<td>Stearylpalmitate</td>
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<td>Cooling 56.54 ± 0.01</td>
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<td>2853.78 ± 0.03</td>
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<td>Palmitoylpealmitate/palmitoyloleate</td>
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<td>2853.98 ± 0.09</td>
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<td>Cooling 45.28 ± 0.09</td>
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<td>Stearylpalmitate/oleyloleate/cholesterylpalmitate</td>
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<td>Cooling 51.31 ± 0.07</td>
<td>2848.46 ± 0.07</td>
<td>2853.61 ± 0.06</td>
<td>184 ± 24</td>
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*Phase transitions for lipid standards were run at least twice with care to obtain data points in the sharp phase-transition temperature region. Replicate runs were combined, and then the phase-transition parameters were calculated.

The phase-transition temperature of the standard waxes measured in this study were linearly related to the degree of saturation, and similar to the linear relationship between saturation and phase-transition temperature measured for human meibum, which is predominantly wax and other natural membranes (Fig. 6). The data were fit to the line with a slope of −308 and a y intercept of 59.1 (r = 0.87).

CH₂ symmetric stretching frequency was 2849.6 cm⁻¹, indicating that the hydrocarbons of cholesterol alone were ordered.

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The properties of the infrared carbonyl band of human meibum were 1736.0 cm⁻¹ and 1732 cm⁻¹ below the Tm, the carbonyl band of cholesterol ester appeared as a central band were observed in the human meibum at 1741 and 1737 cm⁻¹ (Fig. 9D). This is characteristic of crystal field splitting and orthorhombic packing.

In all the wax samples, two CH 2 bending bands were evident in the mixture of SP and OO (Fig. 7A), especially for PO alone (Fig. 7B). The carbonyl bands for PO (Fig. 7B) were better resolved than for the mixture of SP and OO (Fig. 7A). The difference in the phase-transition temperature for the carbonyl band and CH symmetric stretching band in an equimolar mixture of PS and OO was only 0.4°C (Tables 5, 7). Cholesterylpalmitate exhibited a carbonyl band near 1743 cm⁻¹ in the spectra of the mixture cholesterol/palmitate/SP/OO (Fig. 7D). Above the phase-transition temperature the cholesterol ester-carbonyl band was not visible in the SP/OO mixture. (Fig. 7D, top).

The center of the infrared carbonyl band of human meibum was 1736.0 cm⁻¹ below the Tm, at 15°C. It was only 2.2 cm⁻¹ higher than the carbonyl band of cholesterol alone (Fig. 7C). The bandwidth of the carbonyl band peaked dramatically at the phase-transition temperature (Fig. 7C), because at the phase-transition temperature, two carbonyl bands were evident in the mixture of SP and OO (Fig. 7A), especially for PO alone (Fig. 7B). The carbonyl bands for PO (Fig. 7B) were better resolved than for the mixture of SP and OO (Fig. 7A).

The magnitude of the change in the full bandwidth at half height of the carbonyl band on melting was also small for human meibum, relative to that of the standard waxes. The full bandwidth at half height changed from 16.0 cm⁻¹ below the phase-transition temperature to 17.0 cm⁻¹ above the phase-transition temperature. Two shoulders on both sides of the central band were observed in the human meibum at 1741 and 1752 cm⁻¹ (Fig. 8A).

In all the wax samples, two CH 2 bending bands were evident when the hydrocarbon chains were ordered below the phase-transition temperature (Fig. 9A). This is characteristic of crystal field splitting and orthorhombic packing.

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In all the wax samples, two CH 2 bending bands were evident when the hydrocarbon chains were ordered below the phase-transition temperature (Fig. 9A). This is characteristic of crystal field splitting and orthorhombic packing.
The two CH₂ bending bands converged into one band slightly above the phase-transition temperature (Fig. 9A). Two CH₂ bending bands were present in the spectra of cholesteryl palmitate alone at 24°C (Fig. 9Ae) and below the phase-transition temperature when cholesteryl palmitate was present in the mixture SP/OO (Figs. 1A, 9Ac). The characteristics of the CH₂ bending bands for the SP/OO mixture with cholesterol were indistinguishable from those of the mixture of SP/OO and PO without cholesteryl palmitate (Fig. 9A).

Like the CH₂ bending bands discussed above, the two C—C rocking bands confirmed orthorhombic packing below the phase-transition temperature (Fig. 9B). At and above the phase-transition temperature, the two C—C rocking bands converged into one band (Figs. 9B, 9D). Two C—C rocking bands were present in the spectra of cholesteryl palmitate alone at 24°C (Fig. 9C, bottom) and below the phase-transition temperature when cholesteryl palmitate was present in a mixture of SP/OO (Fig. 9C(e)). The characteristics of the C—C rocking bands in the mixture of SP/OO with cholesterol were indistinct.

<table>
<thead>
<tr>
<th>Group</th>
<th>Age/Sex/Ethnicity*</th>
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<td>Group 1, meibum below 14 y</td>
<td>5MC, 4FC, 6MC, 8FC, 10MC, 12FC, 13FC</td>
</tr>
<tr>
<td>Group 2, meibum 14 to 30 y</td>
<td>17MC, 19MC, 19FC, 20FA, 20MA, 21MC, 21MC, 23MC, 23MC, 24MC, 26MA, 32MC</td>
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<tr>
<td>Group 3, meibum above 30 y</td>
<td>52FC, 54MC, 62MC, 65MC, 67MC, 80FB, 85MC, 88MC</td>
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</table>

* M, male; F, female; A, Asian; B, black; C, Caucasian.

**TABLE 6.** Human Meibum Donor Data Used in Figure 6

**FIGURE 7.** Infrared spectra of the carbonyl region for (A) sterylpalmitate/oleylolate, 1:1, m:m. Temperatures (°C) from top to bottom: 70.96, 59.47, 55.45, 52.14, 51.92, 51.37, 50.95, 50.57, 49.41, 49.35, 48.78, 48.02, 47.34, 35.5, and 24.17. (B) Palmityloleate temperatures (°C) from top to bottom: 24.00, 20.25, 20.17, 20.14, 20.09, 20.02, 19.92, 19.85, 19.78, 19.69, 19.63, 19.55, 19.36, 19.16, 18.72, 18.38, 18.35, 17.64, 16.93, 16.80, 15.59, and 12.83. (C) The carbonyl bandwidth at half height (▲) increased then decreased at the transition temperature measured by the carbonyl band center (▲). All data are from heating curves. (D) Infrared spectra of the C=O region: (top) sterylpalmitate/oleylolate, 1:1, m:m, 70.96°C; sterylpalmitate/oleylolate/cholesteryl palmitate, 2:2:1, m:m:m, 70.96°C; sterylpalmitate/oleylolate/cholesteryl palmitate, 2:2:1, m:m:m, 26.07°C; sterylpalmitate/oleylolate, 1:1, m:m, 24.17°C; (bottom) cholesteryl palmitate 24°C.
guishable from the mixture of SP/OO and PO without cholesteryl palmitate (Fig. 15B). The difference between the center of the infrared band \( \text{C} - \text{C} \) rocking bands of human meibum, above and below the phase-transition temperature, was only 1 cm\(^{-1}\) at 729 cm\(^{-1}\) above the phase-transition temperature at 15°C and 721 cm\(^{-1}\) above the phase transition at 55°C. A shoulder at 729 cm\(^{-1}\) was present below the phase-transition temperature (Fig. 8A) that was not as well resolved as the two bands at 730 and 719 cm\(^{-1}\) in the spectra of standard waxes below the phase-transition temperature (Figs. 9C, 9D).

**Discussion**

**Phase 1: Human Meibum Study**

MGD was defined as "a condition of progressive obstruction due to ductal hyperkeratinization or inspissation of secretion." Until this study, inspissation (thickening) of meibum due to ductal hyperkeratinization or inspissation of secretion can now be quantified by infrared spectroscopy. A major finding of this study is that the average order of Md at the physiological temperature, a small 4°C change in phase-transition temperature is enough to cause a 30% increase in lipid disorder so a decrease in double bonds rather than a change in the composition of cholesterol esters is more likely to account for the higher lipid order observed for Md than for Mn.

Compositional changes that could cause Md to become more ordered than Mn include increased hydrocarbon chain lengths and saturation or changes in cholesterol ester content. Using model waxes we found that the change from one double bond to none raised the phase-transition temperature of standard waxes by 30°C (see detailed discussion below). Cholesterol esters caused no change in wax conformation, so a decrease in double bonds rather than a change in the composition of cholesterol esters is more likely to account for the higher lipid order observed for Md than for Mn.

**FIGURE 8.** Average infrared spectra of human meibum showing (A) the carbonyl band region. Top: cholesteryl palmitate; middle: above the Tm at 55°C; bottom: below the Tm at 15°C. (B) The C–C rocking band region. Top: above the Tm at 55°C; middle: below the Tm at 15°C; bottom: cholesteryl palmitate.
Proteins may also influence meibum conformation. Infrared spectroscopy detected significant levels of protein in human meibum before its interaction with tear proteins. In Md, the amount of protein indirectly correlated with lipid order. Studies to directly correlate meibum conformation with the amount of meibum protein are currently being pursued.

Eighteen percent of the patients diagnosed with MGD exhibited meibum lipid phase-transition temperatures that were similar to those for normal donors. Md ordering in these patients could have been transient, coincidental, or associated with other factors, such as ductal hyperkeratinization or inflammation, which contributed to the dry-eye symptoms. Md order at 33.4°C was strongly related to the lipid phase-transition temperature. Factors that contribute to changes in the phase-transition temperature are likely to change Mn and Md in the disordered state at 65°C are not completely disordered as a liquid (melted solid) and are considered liquid crystals. Similarly, Mn andMd in an ordered state at 5°C are not completely ordered as are solids and so we use the term phase-transition temperature rather than melting temperature.

Phase-transition temperatures span 12°C from sample to sample which is much larger than the ±2°C precision of the measurements (Fig. 3A). The variability from sample to sample must be due to lipid compositional differences. The lipid phase-transition temperature is similar but not identical with melting. Mn and Md in the disordered state at 65°C are not completely disordered as a liquid (melted solid) and are considered liquid crystals. Similarly, Mn and Md in an ordered state at 5°C are not completely ordered as are solids and so we use the term phase-transition temperature rather than melting temperature.

The average phase-transition temperature for Md was approximately 4°C higher than that for age-matched Mn. Terada et al. reported the only study that evaluated Md “melting” at 34.0 ± 1.3°C, but this melting temperature was assessed visually. “Melting” of Mn from age-matched controls ranged from 32°C to 37°C in Terada et al. and the SE was too large to be significantly different from the melting of Md. The phase-transition temperature of Mn, 28.5°C (Table 4), was below the
temperature of the eyelid, 33.1°C to 33.4°C, ensuring that the Mn is predominantly disordered at lid temperature. The phase-transition temperature of Md, 32.2°C, is similar to the eyelid temperature, resulting in 47% of the lipid being ordered at lid temperature.

Insight into the functional relevance of a more ordered Md compared with Mn may be gained from two very early studies. Brown and Dervichian found that meibum lipid that are ordered do not spread when placed on the surface of aqueous saline but spread above 35°C, the melting temperature of the meibum. Remarkably, with only a looped piece of thread, they were able to measure the surface tension of meibum lipid layered on saline solution to be 15 dynes/cm, similar to the value measured by modern instruments. Based on these observations, it is reasonable to expect that ordered Md would not spread or perhaps would spread more slowly on the tear film surface even if it were to be expressed from meibomian glands. Like Brown and Dervichian, Holly reported that meibum lipid on saline has "poor spreading characteristics" and "a low film pressure therefore (high surface tension)" The situation in vivo is more complicated than in these in vitro studies. The aqueous tear film contains proteins and surfactants that interact with the meibum lipid. In Holly's seminal paper, he reported that mucin increases the surface pressure of meibum lipid threefold. Recent Langmuir trough experiments have shown that above the phase-transition temperature, the area of the meibum film per molecule is significantly larger. These results provide insight, but one must be cautious in comparing them with ours, since the trough experiments use chloroform, which alters lipid packing and denatures the proteins present in native meibum. As a consequence of Md's having 72% stronger lipid–lipid interactions than Mn, the Md would be expected to have a lower film pressure than Mn and would not spread on the surface of tears as readily as Mn. Brown and Dervichian proposed that meibum lipid disordered by temperature spreads more readily on a saline surface because surface active molecules are free to diffuse and "spread on the watery surface and carry on their back the thick nonpolar hydrocarbon portion of the oil." It is likely that lysozyme, lipocalin, and mucin bind to meibum at the surface, adding stability to the film. The tear proteins lowered the surface tension and increased surface pressure by causing a compression of the lipids. It has yet to be determined how more ordered Md would interact with the tear proteins.

Phase 2: Model Lipid/Wax Study

The characteristics of the lipid phase transitions of human meibum change with age, meibomian gland disease (this study), and therapy. The biophysical changes reflect compositional changes that have been elusive to quantify. For instance, six studies have shown that wax esters compose from 13% to 68% of meibum, hydrocarbons 1% to 38%, sterol esters 8% to 39%, and triglycerides 2% to 43%. Infrared and Raman spectra suggest that wax and sterol esters are disordered in native meibum, so in this study, the phase characteristics of pure waxes and simple mixtures were examined to gain insight on the compositional factors that are likely to contribute to the changes in the phase-transition parameters of human meibum with age, disease, and therapy.

Carron's Region. The carron's band frequency of standard waxes shifted over 7 cm⁻¹ on melting, but the carron's band is not a good indicator of lipid phase transitions for human meibum which shifted by only 2 cm⁻¹. The carron's full bandwidth at half height and frequency are dependent on the environment of the carron's band and not on conformation. When the carbonyl is in an environment with a low dielectric constant such as n-hexane (ε = 1.89) its frequency is 1750 cm⁻¹ with a narrow bandwidth of 12. When the carbonyl is in an environment with a stronger dielectric constant such as CCl₄ (ε = 2.24) or CHCl₃ (ε = 4.8) the carbonyl band frequency is lower, 1742 and 1752 cm⁻¹, and the full bandwidth at half height is higher, 13 and 20 cm⁻¹, respectively. The carbonyl band therefore may be used as an internal probe of the environment surrounding the carbonyl group.

Below the phase-transition temperature, the carbonyl moieties of standard waxes are in an environment similar to CCl₄. On melting, there is a dramatic change to a more hydrophobic environment similar to that of CHCl₃. Waxes are tubelike structures that pack together in multimellar monolayers (Fig. 1). In a system of standard waxes, there are only two environments that the wax carbonyl moiety can experience; the hydrophobic environment of the hydrocarbon chains that have a low dielectric constant and the ester region of adjacent waxes that has a higher dielectric constant. For standard waxes below the phase-transition temperature, the acyl linkages may be in close proximity to the acyl linkages of adjacent waxes. Above the phase-transition temperature, the carbonyl is in proximity to the hydrocarbon region of adjacent waxes.

Below the phase-transition temperature, the carbonyl groups in native human meibum are in an environment with a higher dielectric constant between that of CCl₄ and CHCl₃. Above the phase-transition temperature the environment changes slightly to a more hydrophobic environment with a lower dielectric constant.

Cholesterol Ester. The frequency of the carbonyl of the cholesterylpalmitate ester near 1740 cm⁻¹ is high relative to waxes and meibum, indicating that the ester moiety is in the environment of the hydrocarbon chains of adjacent waxes or the palmitate hydrocarbon region of the cholesterol ester. The small carbonyl shoulder at 1737 cm⁻¹ indicates that a portion of ester moieties are adjacent to the acyl linkages of neighboring wax or cholesterol esters. The CH₂ symmetric stretching band frequency of 1749.6 cm⁻¹ for cholesterylpalmitate indicates that the palmitoyl CH₂ groups are mostly ordered, in trans conformations.

Cholesterol ester had a minimal affect on the phase transition of a mixture of SP and OO. Cholesterol ester at a concentration of 20% increased the Tm of wax without cholesterol ester by only slightly more than 1°C. It is doubtful that smaller changes in cholesterol ester with age and MGD could account for the 4°C shift in the phase-transition temperature of meibum observed with age and MGD (this study).

It would be interesting to study the structure of cholesterol esters in meibum and wax using infrared spectroscopy and deuterated ¹³C isoforms of cholesterol esters. Because cholesterol esters caused little change in the phase-transition temperature of waxes, such a study was deferred.

Phase Transition Temperature. In the present study the amount of hydrocarbon trans and gauche rotomers was used to quantify the phase-transition temperature. A change in phase from an ordered gel phase with more trans rotomers, to a disordered liquid-crystalline phase with more gauche rotomers was accompanied by the disappearance of orthorhombic packing (two bands to one band, Fig 9A). A change in hydrocarbon chain length could cause a change in the phase-transi- tion temperature. In this study an additional CH₂—CH₂ moiety raised the phase-transition temperature approximately 5°C. A similar result using a more extensive group of waxes was found with infrared spectroscopy and a less precise melting point determination. Shorter chain, saturated waxes (myristyl or lauryl stearate) are tilted relative to the lattice plane. Longer chain, saturated waxes (palmitoyl palmitate or palmitoyl stearate) are vertical relative to the lattice plane. Relative to saturated waxes, the addition of one CH₂=CH moiety dramatically low-
tered the phase-transition temperature by approximately 40°C, and the addition of a second CH=CH moiety to a monounsaturated wax decreased the phase-transition temperature by approximately 19°C. The dramatic decrease in phase-transition temperature with unsaturation was expected based on results in previous studies.61

In the present study a unique and unexpected result was that, in mixtures of saturated and unsaturated waxes, the phase-transition temperature of the mixture was disproportionately 30°C to 50°C higher than that of the pure unsaturated waxes, but only 6°C below the phase-transition temperature of pure saturated waxes. The cooperativity of the mixtures was lower than that of pure saturated waxes. Therefore, even though unsaturation has a profound affect on the phase-transition temperature of pure waxes, the effect is much less pronounced in mixtures. In an x-ray diffraction study, a disproportionate difference in the tilt angle of the hydrocarbon chains in mixtures of waxes was noted.55 The hydrocarbon chains of saturated lipids used in the present study, PP and PS, align vertically (A form) with reference to the lattice planes.55 The hydrocarbon chains of lower melting point waxes such as myristylstearate or laurylstearate have a tilted alignment (B form) with reference to the lattice planes.55 When the higher melting A form is mixed with the lower melting B form, the mixture always adopts the B form.55 We believe that in general, the phase-transition temperature of complex mixtures of natural lipids is determined by the level of unsaturation of lipids (Fig. 5A). It is interesting that human lens, muscle and rod outer segment lipids that do not contain waxes fit so closely to the phase-transition, temperature-saturation curve of standard waxes used in this study (Fig. 5A). Patel et al.61 noted that, regardless of whether the lipid was a wax, hydrocarbon, phospholipid, fatty acid, or triglyceride, every carbon atom added to the hydrocarbon chains raised the phase-transition temperature by 1°C to 2°C. The ester linkage decreases the phase-transition temperature by approximately 15°C relative to hydrocarbons containing the same number of carbon atoms.51,75 Changes in lipid saturation and hydrocarbon chain length could account for the 4°C shift in the phase-transition temperature of meibum observed with age49 and meibomian gland dysfunction (this study). Saturation levels of a limited sample of Mn and Md measured using Raman spectroscopy indicates that meibum saturation does not change dramatically with age and meibomian gland dysfunction.41 In the same study, the level of highly unsaturated carotenoids decreased with age and meibomian gland dysfunction.41 An increase in overall chain length by as little as two carbons could increase the phase-transition temperature by 5°C.

The functional relevance of Md’s having a higher phase-transition temperature and order than age-matched Mn, as discussed above, is speculative at this time. Therefore, studies are under way to determine the relationships between meibum lipid composition, structure/conformation, and function. Infrared and Raman spectroscopies are ideal for this purpose, since they both provide conformational and limited compositional information but are nondestructive. It might be informative to spike Md with disordered, unsaturated waxes or Mn with ordered, saturated waxes to determine whether spiked lipids could restore lower lipid order in Md or elevate lipid order in Mn. Nuclear magnetic resonance (NMR) spectroscopy and matrix-assisted mass spectrometry are ideal for quantifying meibum lipid classes and specific composition, respectively. Further studies employing these techniques should elucidate the composition/structure contribution to ordering of the lipids.

References


