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Fatigue Crack Initiation and Propagation in Polyamide-6 and in Polyamide-6 Nanocomposites

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Recent developments in polymer nanocomposites have led to improvements in conventional short-term, but the long-term mechanical properties have received little attention. The objective of the present study was to characterize the effect of nanoparticles on the fatigue crack initiation and propagation mechanisms and on the fatigue properties of polyamide-6 (PA6) nanocomposite (PA6NC) prepared by in situ polymerization with montmorillonite clay. Two approaches were employed: fatigue life measurements and crack growth monitoring. Compared with non-filled PA6 at the same stress amplitude, the number of cycles to fracture was higher for the nanocomposite, which suggests an increase in the intrinsic resistance of the material to crack initiation. However, the crack growth rate results indicated that nanoparticles decreased the resistance to crack propagation. Post-fatigue fractographic observations indicated a change in the fatigue crack propagation mechanism resulting from the addition of nanoparticles, primarily attributed to the increase in yield stress, which favors the development of a fibrillated deformation zone. The fibrillation process in the relatively high crack propagation rate regime appeared to be preceded by plastic deformation at approximately constant volume. Most of the effect of nanoparticles on the fatigue behavior and properties results probably from the mechanical reinforcement on the microstructure and its effect on the yield stress and Young's modulus rather than from the effect of the inorganic filler to act as a stress concentrator. Polym. Compos. 25:433-441, 2004. © 2004 Society of Plastics Engineers.

INTRODUCTION

Inorganic microparticles and microfibers are often added to semicrystalline polymers employed for structural applications. These particles increase the stiffness, and, when the cohesion between them and the matrix is maintained up to high stresses, they also increase the strength (1). However, since particle-reinforced polymers are frequently used in applications where constant or cyclic stresses are applied, the effect of particles on the creep and fatigue properties has to be considered in estimations of the overall mechanical performance. The fatigue performance of polymers can be quantified by two conventional methods: characterization of the fatigue crack initiation resistance on smooth specimens and characterization of the fatigue crack propagation (FCP) resistance employing precracked specimens (2).

During fatigue crack initiation in reinforced semicrystalline polymers, cyclic loading generates strain mismatches at particle-matrix interfaces, which generally results in local debonding. This debonding induces a zone of stress concentration in the matrix, which can

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lead to fatigue crack initiation. When mineral microparticles are added at low concentrations for which the stiffness of polypropylene or polyamide 6 (PA6) is not significantly affected, the resistance to fatigue crack initiation is found to decrease, and this effect becomes more important as the maximum dimensions of the particles increase (3–5). By adding glass fibers at concentrations that significantly increase the stiffness of PA6, the number of cycles to fracture is found to increase at a given stress amplitude and to decrease at a given initial strain amplitude (6–8). The overall effect of inorganic fillers on fatigue crack initiation can be rationalized by a weakening effect attributed to local stress concentrations at particles and by a strengthening effect attributed to an increased stiffness.

Industrial processes have been developed to obtain good dispersion of sheet-like inorganic nanoparticles, approximately 1 nm in thickness and $0.1-1.0 \ \mu m$ in lateral dimensions, within a limited number of semicrystalline polymers (9). Unlike their micro-sized counterparts, these nanoparticles can largely improve the tensile modulus and ultimately the tensile strength without significantly changing the density. The improvements in stiffness can be related to the high aspect ratio of the fully exfoliated nanoparticles and to the level of exfoliation. In addition, the significant increases in tensile strength seen in polyamide-based systems are usually explained by strong particle-matrix interactions (9). These strong interactions can have direct effects on the interface strength, the level of exfoliation, the chain mobility and the supermolecular structure. A decrease in chain mobility by adsorption at the surface of the nanoparticles has been proposed to explain the decrease in free volume fractions (10). Calculations have suggested that crystallization in the presence of nanoparticles becomes a bi-dimensional rather than a tri-dimensional process (11). TEM observations have indicated that nanoparticles tend to orient themselves parallel to the shear flow plane (12, 13) and that crystallites in polyamide-12 become preferentially oriented perpendicularly to that plane (13). These orientation effects can explain the higher increases in tensile properties measured when an in situ polymerized PA6-based nanocomposite is injection molded instead of compression molded (14).

While the effects of clay nanoparticles on the tensile properties of semicrystalline polymers have been well characterized, only few studies have been reported on the effect of nanoparticles on the fatigue performance of nanocomposites. Strain monitoring during fatigue of a PA6-based nanocomposite has previously been reported (15, 16); however, attention was drawn primarily to the thermal instability phenomenon. This phenomenon results from cycling a large volume of material above a critical stress amplitude or loading frequency leading to hysteretic heating (17, 18). In a previous paper (19), the effect of nanoparticles on the evolution of bulk fatigue damage has been reported. It was found that the macroscopic stress-strain-time response evolved in a succession of three regimes associated with different viscoelastic deformation processes. Nanoparticles were found to reduce the deformation rate in the first, but this effect was not sustained in the following regimes. It was proposed that axial fatigue loading in the last regime of viscoelastic deformation resulted in the nucleation and growth of microvoids.

The present paper discusses the effect of clay nanoparticles on the mechanisms of fatigue fracture and on the fatigue performance of a PA6. The fatigue performance was evaluated by generating curves of the stress versus the number of cycles to fracture (S-N curves) on non-notched axial specimens as well as curves of the fatigue crack propagation rate (da/dN) versus the stress intensity factor (ΔK) on pre-cracked specimens. The mechanisms of fatigue fracture were characterized by fractographic observations as well as by post-fatigue observations on a transverse cross section near the FCP plane.

EXPERIMENTAL METHOD

The polymer nanocomposite employed in the present study was obtained from UBE Industries as commercial grade 1015C2. As described elsewhere (15), this grade is produced by in situ polymerization. Montmorillonite, a natural layered crystal member of the 2:1 phyllosilicate family, is first hydrated in the presence of hydrochloric and aminolauric acids in order to replace the sodium cations present in the interlayers by acid functions. Two weight-percent (2 wt%) of this surface-modified product is subsequently mixed with ε -caprolactam monomers before polymerization. The resulting PA6-based nanocomposite (PA6NC) contains exfoliated nanoparticles with approximate thickness and lateral dimensions of 1 nm and 100 nm, respectively (12). The non-filled PA6 employed in the present study, obtained from UBE Industries as commercial grade 1015B, had the same molecular weight ($\overline{M}_n = 22,200$ g/mol) as the PA6NC (20).

All specimens were prepared following the same procedure. Pellets of PA6NC and PA6 were dried for 2 days at 80°C under vacuum. Each material was then injection molded with an increasing melt profile temperature ranging from 230°C to 250°C and a mold temperature of 80°C. Prior to mechanical testing, all specimens were further dried for two weeks at 80°C under vacuum in order to remove any trace of humidity that can result from processing because of the hygroscopic nature of PA6. The glass transition temperature, T_g , was then determined from loss shear modulus vs. temperature curves, obtained employing a rheometric mechanical spectrometer at a frequency of 1 Hz and at a constant rate of temperature increase of 2° C/min. A similar T_a of approximately 65°C was obtained for both materials. Tensile test results reported elsewhere (21), on specimens processed and conditioned following the procedure presently described, indicated room temperature values of Young's modulus, ultimate tensile strength (UTS), and strain at fracture respectively of 4.1 GPa, 100 MPa, and less than 10% for PA6NC and respectively of 2.7 GPa, 74 MPa and 175% for PA6.

The non-notched axial fatigue specimens were molded directly to their final dimensions. Following the general recommendations of ASTM E606, these specimens had a nominal thickness of 3.1 mm and a gauge section width and length of 10 mm and 14 mm, respectively. The radius of curvature at the shoulders was 15 mm. Load-controlled axial fatigue tests were carried out in tension-tension using servo-hydraulic testing machines (Instron 8511 and MTS 415). For all tests, the load or R-ratio (R = $\sigma_{min}/\sigma_{max}$) was maintained constant at 0.1, the cyclic waveform was sinusoidal and the cycling frequency was 5 Hz. During a few experiments, the specimen temperature was monitored using a K-type thermocouple attached to the surface. All axial fatigue and fatigue crack propagation tests were carried out at room temperature.

The fatigue crack propagation tests were done following the ASTM E-647 test method. Compact tension specimens were machined from injection-molded plates to a width (W) of 50 mm, a notch depth of 11 mm and a notch opening width of 2 mm. The specimen thickness was kept equal to the plate thickness of 5 mm; the notch and cracking plane was perpendicular to the direction of injection flow. Tests were performed employing an Instron 8511 servo-hydraulic testing machine at an R-ratio of 0.1 and with a sinusoidal waveform. A razor-blade cut was introduced manually at the notch tip to produce a starter crack just before fatigue pre-cracking, which was carried out at a frequency of 20 Hz and over a distance of 2 mm. The crack length was monitored online using the compliance measured from a clip-on COD gauge. From this crack length and from the imposed load, the amplitude of the variation in the stress intensity factor (ΔK) was computed. The initial ΔK value was set equal to or slightly higher than the ΔK applied during pre-cracking; the ΔK gradient ranged from 0.04 to 0.14 MPa.m^{1/2}/mm but was constant during all tests. Each crack growth rate measurement was obtained over a propagation distance of 0.15 mm. In all tests, final fracture occurred before the crack length reached 30 mm, thus ensuring the applicability of linear elastic fracture mechanics.

The fracture surfaces of several specimens were observed after the fatigue tests. These fractographic observations were carried out with a JEOL JSM-6100 scanning electron microscope (SEM), after depositing a palladium-gold layer on the fracture surfaces by ion sputtering. Also, a section perpendicular to the crack plane was prepared by ultra-cryomicrotomy, coated with palladium and observed with an Hitachi S4700 field emission gun scanning electron microscope (FEGSEM).

RESULTS AND DISCUSSION

Axial Fatigue

Quantitative results from the axial fatigue tests are presented in *Fig. 1*, where the fatigue life (*N*) measured is plotted as a function of the imposed constant maximum cyclic stress (σ_{max}). For PA6NC, the response can



Fig. 1. Maximum cyclic stress versus fatigue life: (■) PA6NC, thermal regime; (□) PA6NC, mechanical regime; (●) PA6, thermal regime; (○) PA6, mechanical regime.

be divided into two regimes. In the high-stress regime, approximately above 78 MPa or 78% of the UTS, fracture occurred in a macroscopically ductile manner by necking. Just below this high-stress regime, at a maximum cyclic stress of 75 MPa, the average initial strain amplitude was measured to be 0.98%. For this maximum cyclic stress of 75 MPa, the specimens failed without any necking. Figure 1 shows that the dependence of N on σ_{max} is more important in the higher stress regime. The macroscopic fracture process for PA6 also changed from macroscopically ductile to macroscopically brittle at a given critical $\sigma_{max},$ which was found to be approximately 60 MPa or 81% of the UTS. The initial strain amplitude at a slightly lower σ_{max} of 57 MPa, was 0.93%. This critical σ_{max} value was employed to divide the response of PA6 into low-stress and high-stress regimes, as this has been previously reported for several other similar material systems (2, 18, 22). Although the critical σ_{max} is higher for PA6NC than for PA6, the relative stress value with respect to the UTS and also the initial strain amplitude at which this transition occurs are very similar.

In the high-stress regime, the temperature rose just before necking, suggesting the occurrence of the wellknown thermal instability phenomenon (2, 18), caused by poor dissipation of the heat generated by viscoelastic damping. However, experiments employing a trapezoidal cyclic waveform with varying hold times at maximum and minimum stresses showed that the thermal instability was controlled by a threshold value of the accumulated strain in the specimen rather than by the testing frequency (23). Therefore, necking could also be caused by a mechanical instability resulting from a competition between strain hardening and geometrical softening, which is the general origin of necking during tensile elongation tests (24). Independently of its thermal or mechanical origin, the instability leading to necking must be regarded as an artifact of the loading mode.

In the lower stress regime, detailed fractographic observations (see the Fatigue Damage Characterization section) provided evidence that sub-critical fatigue crack propagation occurred before the final macroscopically brittle fracture. The fatigue life in that regime is thus more representative of the overall resistance of the specimens to the initiation and propagation of fatigue cracks. From the results presented in Fig. 1, this overall mechanical fatigue resistance of PA6NC and PA6 can be compared by employing the results obtained for each of the three following conditions: PA6NC at 57 MPa, PA6NC at 75 MPa, and PA6 at 57 MPa. As shown in Table 1, the average values of the initial strain amplitudes for PA6NC at 75 MPa and PA6 at 57 MPa are similar. Table 1 presents the average fatigue life for each of the three conditions. When compared to PA6, the average fatigue life of the non-notched PA6NC specimens is more than twice as high at the same σ_{max} value of 57 MPa, but more than three times lower at a similar initial strain amplitude of approximately 0.9–1.0%. Since the curves in Fig. 1 reflect the initiation and propagation of fatigue cracks, analysis of the causes of these differences requires consideration of the fatigue crack propagation tests and of the fractographic observations.

Fatigue Crack Propagation

The macroscopic fatigue crack propagation rate (da/dN) was monitored as a function of the range of stress intensity factor (ΔK) on pre-cracked specimens.

Of the eight fatigue crack propagation tests performed, two were deemed to be invalid. Both tests were on PA6NC with an initial ΔK of 0.8 MPa.m^{1/2} and a ΔK gradient of 0.04 MPa.m^{1/2}/mm⁻¹ and both presented a long plateau, where da/dN remained constant as the macroscopic ΔK increased. This occurred for crack lengths where the fracture surfaces presented several curved crack propagation lines oriented at high angles to the macroscopic crack growth direction (23). Long crack arrest lines approximately perpendicular to the curved crack propagation lines were also observed for these two tests. These observations indicated an important curvature of the crack front. The occurrence of a long propagation plateau in these two tests was considered to result from the increase in the effective

Table 1. Average Fatigue Life (Average Made on the Logarithm of the Fatigue Life) for PA6NC and PA6 Tested at the Same Maximum Stress and at a Similar Initial Strain Amplitude.

Material	PA6NC		PA6
Maximum cyclic stress ($\sigma_{\rm max}$)	57 MPa	75 MPa	57 MPa
Average initial strain amplitude (%)	0.69	0.98	0.93
Average fatigue life (kilocycles)	5300	600	2300

length of the crack front artificially increasing the resistance to fatigue crack growth by increasing the amount of deformed material available to sustain the applied load. One or two smaller plateaus were observed during each of the two other fatigue crack propagation tests performed on PA6NC with an initial ΔK of 0.45 MPa.m^{1/2}.

The results considered to be valid obtained from the crack propagation tests are plotted on a log-log scale in *Fig. 2*. The crack growth rate at any given ΔK is comparatively higher in PA6NC than in PA6. As shown in *Fig. 2*, a linear log-log relationship between da/dN and ΔK was observed during the test on PA6NC at an initial ΔK of 0.8 MPa.m^{1/2} as well as during all three tests on PA6. This linearity is also generally found in a large variety of polymeric systems (2, 25, 26), including glass-fiber reinforced PA 6 (27, 28).

Any deviation from linearity can be attributed to experimental artifacts. For the two tests on PA6NC at an initial ΔK of 0.45 MPa.m^{1/2} and a ΔK gradient of 0.14 mm^{-1} , the small plateaus can be attributed to crack curvature effects, considerably less important than but of the same nature as those which caused two tests to be classified as invalid. The high susceptibility for crack curvature effects during the present tests on PA6NC can be rationalized from microstructural gradients and from the high sensitivity of the cracking mechanism to hydrostatic stresses (see the General Discussion section). This crack curvature leads to an overestimation of the intrinsic FCP resistance. Since the intrinsic resistance of the materials to fatigue crack propagation should be represented by the linear portions (neglecting any plateaus) of the log-log plots, Fig. 2 indicates that this resistance is significantly lower in PA6NC than in PA6.



Fig. 2. Macroscopic fatigue crack propagation rate versus range of stress intensity factor: $(\bigtriangledown, \bigcirc)$ PA6NC, initial $\Delta K = 0.45$ MPa.m^{1/2}, ΔK gradient = 0.14 MPa.m^{1/2}/mm; (\Box) PA6NC, initial $\Delta K = 0.8$ MPa.m^{1/2}, ΔK gradient = 0.04 MPa.m^{1/2}/mm; (\blacksquare) PA6, initial $\Delta K = 0.6$ MPa.m^{1/2}, ΔK gradient = 0.04 MPa.m^{1/2}/mm; (\blacksquare) PA6, initial $\Delta K = 0.9$ MPa.m^{1/2}, ΔK gradient = 0.04 MPa.m^{1/2}/mm; (\blacksquare) PA6, initial $\Delta K = 0.9$ MPa.m^{1/2}, ΔK gradient = 0.04 MPa.m^{1/2}/mm.

Fatigue Damage Characterization

The fracture surfaces were observed by SEM. *Figure 3* presents a typical example of the fracture surfaces from the axial tests, which surfaces all contained two distinct regions: a relatively small flat central region approximately perpendicular to the load axis and a surrounding region inclined very approximately at 45° to this axis. The inclined fracture surface presented numerous quite large crack propagation or ridge lines but otherwise was microscopically featureless. In the flat central region, small crack propagation lines radiated away from the crack initiation site (such as at I in *Fig. 3*). These fractographic features indicate that the flat central region corresponds to the fracture surface produced by fatigue cracking and that the inclined region corresponds to that of final fracture.

A typical fatigue crack initiation site in a PA6NC specimen tested at σ_{max} of 57 MPa is shown in Fig. 4. For all tested specimens, several crack propagation lines emerge from the periphery of what appears to be a micro-heterogeneity. The number of distinct propagation lines indicates that several microscopic cracks initiated simultaneously, which suggests a relatively long crack initiation period from the interface of this heterogeneity. X-ray energy dispersive spectroscopy indicated that the micro-heterogeneities contained variable proportions of silicon, magnesium, calcium and/or aluminum, which suggests that these heterogeneities are inherent clay by-products present in the nanocomposite. Since such a micro-particle was observed at the fatigue initiation site in all PA6NC axial specimens, the presence of such particles clearly promotes fatigue initiation. It thus appears that the axial fatigue test results are inherently affected by the limited level of purity of commercial nanocomposites produced from natural clay minerals.



Fig. 3. Typical fracture surface of an axial fatigue specimen of PA6NC ($\sigma_{max} = 75$ MPa), with a macroscopically flat central region and a surrounding fast fracture region. The tails of the arrows begin on the start of the zone of fast fracture. The site labeled I indicates the fatigue crack initiation region. The sites A and B are references for Figs. 6 and 7.



Fig. 4. Crack propagation lines emerging from the periphery of an heterogeneity at a typical fatigue crack initiation site located in the flat central region of the fracture surface of a PA6NC axial fatigue specimen ($\sigma_{max} = 57$ MPa, fatigue life = 3197 kilocycles).

The evolution of the fractographic details, in moving away from the fatigue initiation site in a PA6NC specimen tested at σ_{max} of 75 MPa, is shown in *Figs. 5, 6* and 7. Very close to the crack initiation site (*Fig. 5*), the fracture surface contains an array of small voids, which are separated by thin polymer ligaments or fibrils. The presence of these fibrils or ligaments indicates that voids formed a fibrillated deformation zone in the material well before complete surface separation. The term "fibrillated deformation zone" is employed to emphasize that the nucleation and growth mechanism of this voided structure is different from that prevailing during crazing in amorphous polymers as well as during the formation of low temperature crazes in semicrystalline polymers (29–32).

As the distance from the crack initiation site increased, the size and length of the fibrils became progressively larger. A typical fibril structure observed at approximately half the distance between the crack initiation site and the end of the flat central region of Fig. 3 (site A) is shown in Fig. 6. As shown in Fig. 7, the length of the fibrils starts to progressively decrease near the end of the flat central region (site B). The surface features observed near the middle of Fig. 7 also show the characteristic structure generally referred to as "island type." It has been proposed (33, 34) that this island type structure results from the crack growing alternatively along the upper and lower interfaces between the fibrillated deformation zone and the surrounding bulk material. This structure is observed in polymers at the end of large planar voided region close to the transition with the final fracture zone.

The fracture surfaces generated during the fatigue propagation tests in PA6NC were also observed. A lamellar fibril structure, observed in the center of the FCP surfaces at a crack length for which the macroscopic



Fig. 5. Array of small voids or dimples separated by thin ligaments in the flat central region and near the fatigue crack initiation site I (Fig. 3) of a PA6NC axial fatigue specimen ($\sigma_{max} = 75$ MPa).



Fig. 7. Decrease in the length of the ligaments near the end of the flat central region at site B (Fig. 3). The crack propagation direction is approximately from left to right. The structure near the middle of this fractograph is referred to as "island type."



Fig. 6. Larger ligaments at site A (Fig. 3) between the crack initiation site and the end of flat central region. The crack propagation direction is approximately from the lower left to the top right corner. The direction normal to the injection plane is horizontal.

da/dN was approximately 10^{-4} mm/cycle, is shown in *Fig. 8.* The morphology of these fibrils is almost identical to that presented in *Fig. 6* and observed on the flat central region of the fracture surface of the axial fatigue specimens. This similarity confirms that fatigue crack propagation and not simply a monotonic fracture after an incubation period occurred in the flat central region of the axial specimens. Large lamellar fibrils such as those observed in *Figs. 6* and *8* appeared to be preferentially oriented perpendicularly to the specimen thickness direction rather than to the macroscopic crack propagation direction.

To further characterize this fatigue crack propagation process, *Fig. 9* presents observations made on a



Fig. 8. Lamellar ligaments observed on the fracture surface of a fatigue crack propagation specimen of PA6NC at a macroscopic da/dN $\approx 10^{-4}$ mm/cycle. These ligaments are similar to those presented in Fig. 6. The macroscopically imposed crack propagation direction is from bottom to top. The direction normal to the injection plane is horizontal.

transverse cross section taken below the fracture surface in the area where *Fig. 8* was taken. A layer of "textured" material having an approximately constant thickness of 3 μ m over a distance of at least 0.2 mm is present just below the fracture surface. The term textured is employed here to account for the presence of small lines (which are vertical in *Fig. 9*) preferentially oriented parallel to the fracture plane. The interface between this textured layer and the bulk material, and the bulk material itself, both contain microvoids, which are clearly opened. The proportion of the surface that corresponds to microvoids is large near the interface and progressively decreases within the first 5–10 μ m



(a)



(b)

Fig. 9. Transverse cross-section showing the structure of a PA6NC FCP specimen just below the crack plane (macroscopic da/dN $\approx 10^{-4}$ mm/cycle). The normal to the crack propagation plane is horizontal and the crack propagation direction is vertical. (a) From left to right: trace of the fracture surface, layer of textured material, interface and bulk material. (b) Higher magnification view centered on the interface.

of bulk material. In contrast, the surface density of similar microvoids appears relatively low in the textured layer. Neither microvoids nor textured layers were observed at edges not corresponding to a fracture surface.

Based on the assumption that the narrow preferentially oriented lines seen in the textured layer correspond to elongated microvoids almost completely closed during deformation ahead of the crack tip, this textured layer appears to consist of material, which has been stretched along the crack propagation direction. Stretching along this direction has been previously reported for monotonic loading (35) of rubber-toughened polyamide 66, with the deformation estimated from the change in the aspect ratio of the pores generated at the rubber microparticles. Although some observations suggest that stretching within the textured layer may be higher near the crack plane, no clear change in the morphology of the stretched polymer within this layer was observed as a function of the distance from this plane.

To rationalize the observations reported here, it is proposed that crack growth was preceded by the advance of an interface, between a highly deformed zone and the bulk material. This process would present similarities with the known mechanism of growth of shear bands in amorphous polymers (36). The presence of a gradient of surface density of microvoids within the bulk material suggest an early stage of localized void formation, which can be expected to promote the initiation and growth of a fibrillated deformation zone near the mid-plane of the highly deformed zone.

To compare the crack propagation mechanisms in PA6NC to that of the non-filled matrix, the fracture surfaces of the axial fatigue PA6 specimens were also observed. These specimens also presented a flat central region, within which the fatigue crack initiated at heterogeneities. The fractographic features of these heterogeneities, however, were different from those observed for PA6NC. A typical crack initiation site in a PA6 specimen tested at σ_{max} of 57 MPa is shown in Fig. 10a, with part of the heterogeneity at which the crack initiated being visible. The heterogeneities found at the crack initiation site in the PA6 specimens presented a relatively smooth morphology. X-ray microanalyses never revealed the presence of inorganic elements in these heterogeneities. A few spherulites (approximately $10 \ \mu m$ in size) are also seen in Fig. 10a, with a typical transspherulitic fracture aspect (37). At an increasing distance from the crack initiation site, the plastic zone size and the crack tip blunting increased as a result of the increase in ΔK and in the crack opening displacement, which led to smoother fractographic features. Figure 10b presents a transitory aspect where the surface is smooth but where the effect of the spherulites in reinitiating the crack can still be resolved clearly. Both near the fatigue initiation site and in an intermediate portion of the fatigue crack propagation zone, the fracture surfaces of the PA6 specimens did not present fibrils similar to those observed in PA6NC, indicating a lower tendency of the material to form microvoids ahead of the fatigue crack tip for these loading conditions.

GENERAL DISCUSSION

Microvoid formation in polymers is time-dependent and stress-dependent and is particularly affected by the hydrostatic component of stress (33, 38). Because of the difference in yield stress, significantly higher stresses ahead of the crack tip are expected in PA6NC than in PA6, which could be the primary reason for the differences in the fatigue cracking mechanisms indicated in these two materials by the fractographic observations. The importance of microvoid formation in the cracking mechanism of PA6NC can explain its higher tendency to undergo crack curvature during the fatigue crack propagation tests. During these tests, the hydrostatic component of the stress in the zone ahead





Fig. 10. Fracture surfaces of axial fatigue specimens of PA6 ($\sigma_{max} = 57$ MPa). (a) Interface (oriented horizontally) with the heterogeneity (bottom) that caused crack initiation and a few spherulites (approximately 10 µm in size) with typical transspherulitic fracture aspect. (b) Large half dimples at approximately half the distance between the crack initiation site and the end of the flat central region. The crack propagation direction is approximately from bottom to top.

of the crack tip is higher near the specimen mid-thickness than near the lateral surfaces. Therefore, voids will nucleate more easily near the mid-thickness, decreasing the local resistance to fatigue propagation.

The different morphology of the fibril arrays observed on the fracture surfaces of PA6NC suggests an effect of the nanoparticles on the cracking mechanism. At low propagation rates (*Fig. 5*), the size of the microvoids between fibrils is of the order of 1 μ m, which is comparable to the maximum expected length of the nanoparticles. However, very similar fracture surface morphologies have been reported for fatigue of nonfilled semicrystalline polymers (39, 40). Thus, while the presence of the nanoparticles may well facilitate the initiation of microvoids especially in the presence of the higher local cyclic stresses expected, the fractographic observations do not allow us to identify this unambiguously.

At higher propagation rates, the lamellar fibrils oriented preferentially perpendicularly to the injection plane could result from preferential orientation of the nanoparticles or of the polymer chains, but similar walled structures have been observed in compressionmolded specimens of non-filled polyethylene (41, 42).

Surface separation leading to crack advance in PA6NC occurs by the breakdown of a fibrillated deformation zone. The fibrillation process requires the nucleation of microvoids in the material, which is easily activated because of the high stresses as well as, possibly, the presence of stress concentrators in this material. Pre-existing voids, crystallite ends and amorphous regions and exfoliated nanoparticles are all possible stress concentrators. It has not been established clearly that the nanoparticles contribute directly to the decrease of the resistance of the PA6 matrix to fatigue crack propagation; however, the indirect effect of the nanoparticles to increase the yield stress, thus facilitating the formation of microvoids ahead of the crack tip, is expected to be an important factor contributing to this decrease in fatigue propagation resistance.

Although the resistance to fatigue crack propagation of PA6 is decreased by the addition of nanoparticles, the fatigue life of axial specimens was increased when the comparison was made for the same stress amplitude. The fatigue life represents the sum of the time spent in initiating the crack and in propagating it to a critical length, which depends on the fracture toughness of the material. Since the fatigue crack propagation resistance and the fracture toughness (21) for PA6NC are lower than for PA6, this increase in fatigue life due to the addition of nanoparticles can be interpreted as an increase in the resistance of the specimens to the initiation of a fatigue crack.

Since all specimens were processed and tested according to the same procedure, this increase in the fatigue resistance of the axial specimens is expected to originate from an actual increase in the fatigue initiation resistance of the material. The decrease in strain amplitude, which results from the increased yield stress, should be a factor contributing to the increase in fatigue resistance of the axial specimens. In a related study (19), nanoparticles were found to decrease the rate of accumulation of viscoelastic deformation in the specimens during the initial stages of the axial fatigue tests, which may also contribute to this increase in fatigue resistance.

CONCLUSIONS

From this study, it can be concluded that:

1. Dispersing sheet-like inorganic nanoparticles in a PA6 matrix increases the fatigue life of axial fatigue specimens subjected to a given tension-tension stress amplitude, but decreases the fatigue life at given strain amplitude. Therefore, the effect of nanoparticles on the fatigue life appears to be analogous to the one reported for inorganic fillers having dimensions in the microscale.

- 2. The increase in fatigue life at given stress amplitude is expected to be a consequence of an increase in the intrinsic material resistance to crack initiation, which is favored by the effect of nanoparticles to increase Young's modulus and accordingly to decrease the displacement amplitude of the macromolecules during cycling.
- 3. Although the addition of nanoparticles results in a change in the cracking mechanism, observations strongly suggested that the development of a fibrillated deformation zone in PA6NC is preceded by the permanent deformation of a relatively thick layer of material.
- 4. The resistance to fatigue crack propagation resistance decreases, which is probably attributed to an indirect effect of the nanoparticles to increase the yield stress that favors the formation of microvoids ahead of the crack tip.
- 5. Most of the effect of nanoparticles on the fatigue behavior and properties results probably from the mechanical reinforcement of the microstructure and from the increase in yield stress and Young's modulus rather than from the effect of the inorganic filler to provide stress concentrators.

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