Accounting for biological and physical sources of acoustic backscatter improves estimates of zooplankton biomass

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Abstract: To convert measurements of backscattered acoustic energy to estimates of abundance and taxonomic information about the zooplankton community, all of the scattering processes in the water column need to be identified and their scattering contributions quantified. Zooplankton populations in the eastern edge of Wilkinson Basin in the Gulf of Maine in the Northwest Atlantic were surveyed in October 1997. Net tow samples at different depths, temperature and salinity profiles, and multiple frequency acoustic backscatter measurements from the upper 200 m of the water column were collected. Zooplankton samples were identified, enumerated, and measured. Temperature and salinity profiles were used to estimate the amount of turbulent microstructure in the water column. These data sets were used with theoretical acoustic scattering models to calculate the contributions of both biological and physical scatterers to the overall measured scattering level. The output of these predictions shows that the dominant source of acoustic backscatter varies with depth and acoustic frequency in this region. By quantifying the contributions from multiple scattering sources, acoustic backscatter becomes a better measure of net-collected zooplankton biomass.

Résumé: Afin de pouvoir convertir des mesures de rétrodiffusion d'énergie acoustique en estimations de l'abondance et en renseignements taxonomiques relatifs à la communauté zooplanctonique, il est nécessaire d'identifier tous les processus de rétrodiffusion en activité dans la colonne d'eau et de mesurer leurs contributions à la rétrodiffusion. Nous avons inventorié les populations de zooplancton sur les bords est du bassin de Wilkinson du golfe du Maine dans le nord-ouest de l'Atlantique en octobre 1997. Nous avons obtenu des échantillons au filet traînant à différentes profondeurs, des profils de température et de salinité, ainsi que des mesures de rétrodiffusion acoustique à des fréquences multiples dans les 200 m supérieurs de la colonne d'eau. Nous avons identifié, dénombré et mesuré les échantillons de zooplancton. Les profils de température et de salinité ont servi à estimer l'importance de la microstructure de la turbulence dans la colonne d'eau. Ces ensembles de données traités avec des modèles théoriques de diffusion acoustique ont permis de calculer les contributions respectives des agents biologiques et physiques de diffusion au niveau global de diffusion mesuré. Le résultat de ces prédictions montre que la source dominante de rétrodiffusion acoustique dans cette région varie en fonction de la profondeur et de la fréquence acoustique. Une fois calculées les contributions des sources multiples de diffusion, la rétrodiffusion acoustique permet de mieux mesurer la biomasse du zooplancton récolté au filet.

[Traduit par la Rédaction]

Introduction

Acoustic surveys of zooplankton and fish offer many advantages over other sampling techniques (Holliday and Pieper 1995; Foote and Stanton 2000). Measurements of acoustic backscatter using a scientific echosounder can be made from ships, buoys, or moorings and thus provide greater spatial coverage or longer time series than conventional sampling techniques such as net tows or diver observations. Although video sampling methods (Davis et al. 1996; Benfield et al. 2001, 2003) provide some of the same advantages as acoustics, they are not used as extensively as

acoustic surveys and typically have much smaller sampling volumes.

One of the difficulties in using acoustic backscatter to measure marine life in the ocean is that the data collected are indirect measures of biota. If aggregations of animals in a region are monospecific and of similar size, theoretical backscatter models can be used to estimate their distribution and abundance (Hewitt and Demer 2000). If more than one frequency of sound is used, then more categories and size classes of animals may be distinguished and their abundance estimated (Martin et al. 1996; Brierley et al. 1998). However, these approaches require that the type of the scatterers

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present in the water column is known. Because of this, zooplankton samples are typically collected by net tows during an acoustic survey. These samples provide taxonomic and size information that can be used to predict the level of acoustic backscatter in the water column.

This prediction is often referred to as the forward problem (FP). For a particular type of backscatterer (usually delineated by either size or taxonomy), the scattering contribution can be found by multiplying the numerical density of scatterers (N_i , in m⁻³) present and the backscatter cross section for an individual scatterer of this type (σ_i , in m²). The backscatter cross section is a function of several parameters, including the size, shape, composition, and orientation of the scatterer and acoustic frequency. The output of the FP is the volume backscatter coefficient (s_v , in m⁻¹), which is found by summing the contributions from the different types of scatterers present

$$(1) s_{\rm v} = \sum_{i} (N_i \sigma_i)$$

These calculated levels of backscatter are then compared with measured values from field surveys. Often, echosounders record the volume backscatter strength (S_v , in dB), which is related to s_v by

(2)
$$S_{\rm v} = 10\log_{10}(s_{\rm v})$$

If the measured and predicted values agree, the inverse problem (IP) (using measured scattering values and theoretical scattering models to determine the number, size, or type of animals present) is more likely to be solved correctly. However, the predicted and measured volume backscatter strengths occasionally differ by an order of magnitude (or more), which can lead to large errors or uncertainty in solving the IP. For example, a 3 dB difference in S_v corresponds to a factor of two difference in the number of animals or their biomass. In practice, solutions to the FP are used in a diagnostic sense to determine how well the theoretical scattering models predict observed levels of backscatter in the ocean (Wiebe et al. 1996, 1997). These results often indicate that only a small subset of the animals present in the water column are acoustically important or, in some cases, that backscatter predictions based on the sampled zooplankton are unable to account for all of the observed backscatter and that another scattering source is unaccounted for in the FP analysis (Mair et al. 2005).

Although nearly every acoustic survey relies on net tow data to ground-truth the acoustic data, many do not take complete advantage of all the available information provided by the net tow. Typically, FP calculations are performed to identify the acoustically dominant taxa in the water column; however, the quantification of backscatter contributions from all of the taxa found in the water column is rarely done. Although many taxa present in the water column will contribute negligibly to the overall level of backscatter (because of small size, low numerical density, or low scattering efficiency), there are often several scattering sources that contribute substantially to the overall level of measured acoustic backscatter. Furthermore, the vertical distribution of zooplankton in the water column varies, which may cause the taxa that is the largest acoustic scatterer to change as a

function of depth. Many of the net tows used to ground-truth acoustic data cannot provide this information, which may further complicate estimates of zooplankton abundance or distribution.

Even if the backscatter from marine organisms is accurately measured, there are other processes in the ocean that can contribute measurable amounts of backscattered acoustic energy. Suspended sediments, air bubbles, ocean mixing processes, and other biota have all been observed during acoustic surveys of zooplankton (Wiebe et al. 1997; Trevorrow 1998); however, backscatter from nonbiological processes is rarely quantified during field surveys.

Acoustic methods have been used for many years to observe physical mixing processes in the ocean (Thorpe and Brubaker 1983; Orr et al. 2000; Ross and Lueck 2005). It has only been in the last decade that theoretical scattering models for these processes have begun to be tested in the field. These models use the variations in temperature and salinity to calculate changes in the index of refraction and density in the water column that result from turbulence and other mixing processes. The acoustic scattering that occurs from these variations can then be predicted (Goodman 1990; Seim et al. 1995; Lavery et al. 2003).

Depending on the mixing rates present (generally characterized by the dissipation rates of turbulent kinetic energy (ϵ) and temperature variance (χ)), backscatter from turbulent microstructure can be equal to or greater than that from assemblages of zooplankton, particularly at the lower range of frequencies commonly used for acoustic surveys (i.e., <100 kHz; see fig. 1 in Warren et al. 2003). Estimates of S_v from this mechanism range from -110 dB in calm waters with small temperature and salinity stratification to -60 dB or higher for regions of intense mixing such as the Bosporus Strait (Seim 1999).

This study examines the contributions of both biological and physical sources of backscatter to the water column in the Gulf of Maine. Contributions from each scattering source were quantified using theoretical scattering models and either net tow data or hydrographic profiles for multiple depth bins of the water column and multiple acoustic frequencies. The theoretical predictions of backscatter from the different scattering sources were used to correct the amount of measured backscatter in the water column to reflect scattering only from biological sources. The adjusted values of measured backscatter were then compared with measurements of zooplankton biomass.

Materials and methods

As part of a GLOBEC (GLOBal ocean ECosystem dynamics) process cruise studying the populations of *Calanus* in the basins of the Gulf of Maine, an acoustic survey was conducted in the eastern part of Wilkinson Basin (located between Georges Bank and Stellwagen Bank) in mid-October 1997 from the R/V *Endeavor* (Table 1). To provide spectral information about the acoustic backscatter processes occurring in the water column, multiple frequency acoustic backscatter data were collected by BIOMAPER-II (BIO-optical Multifrequency Acoustical and Physical Environmental Recorder) (Wiebe et al. 2002).

Multiple opening and closing net and environmental

Table 1. Location and time of the CTD vertical profiles and MOCNESS tows used in this study collected in October 1997 from the R/V *Endeavor*.

Event	Julian year-day	Latitude (N)	Longitude (W)	Beginning or end
CTD 08	287.451	42°14.97′	68°44.77′	Beginning
	287.467	$42^{\circ}14.97^{\prime}$	68°44.77′	End
MOC 07	287.620	$42^{\circ}24.04^{\prime}$	68°49.03′	Beginning
	287.686	42°24.93′	68°44.22′	End
CTD 10	289.535	$42^{\circ}25.08'$	68°44.49′	Beginning
	289.562	42°25.08′	68°44.49′	End
MOC 09	289.896	$42^{\circ}28.70'$	68°45.00′	Beginning
	289.949	$42^{\circ}30.97^{\prime}$	68°46.69′	End

sensing system (MOCNESS) tows (Wiebe et al. 1985) were conducted to collect zooplankton samples while BIOMAPER-II was concurrently recording acoustic backscatter data from the water column. Profiles of the temperature and salinity of the water column were made with conductivitytemperature-depth (CTD) sensors onboard the MOCNESS and nearby higher-resolution CTD casts (Sea-Bird 911; Sea-Bird Electronics, Inc., Bellevue, Wash.) from the vessel. Data from two sampling periods are presented herein, with samples from year-day 287 (CTD 08, MOC 07) and year-day 289 (CTD 10, MOC 09). The CTD and MOC-NESS sampling took place in the same general area (Table 1). These data sets (acoustic backscatter, zooplankton taxa and size, and temperature and salinity profiles) provided enough information to estimate the contributions from biological and physical sources of acoustic backscatter.

Acoustic backscatter measurements

BIOMAPER-II (Wiebe et al. 2002) is a towed body with numerous acoustic, environmental, video, and bio-optical sensors. The acoustic system consists of five pairs of transducers (operating at 43, 120, 200, 420, and 1000 kHz), with one of each frequency looking upward and the other downward. The transducers have depth ranges of 200, 200, 150, 100, and 35 m, respectively, with a vertical resolution of 1 m depth bins. Backscattered energy from each transducer and for each depth bin was recorded as echo-integrated volume backscattering strength every 12 s. At typical survey speeds, this ping rate corresponds to a horizontal range between pings of 20 to 50 m. The instrument was normally towed obliquely through the water column; however, because additional equipment was in the water during these measurements, the tow body was kept at a constant depth of approximately 5 m below the surface, and slower tow speeds resulted in a horizontal resolution between pings of approximately 10 to 15 m. Because of this configuration and the limited depth range of the 1 MHz transducer, only data from the downward-looking transducers at 43, 120, 200, and 420 kHz were analyzed.

The acoustic data were processed and combined with data from the ESS (environmental sensing system) sensors that are also onboard BIOMAPER-II. The acoustic system was calibrated using standard target spheres before the cruise. The final data file provided echo-integrated volume back-scatter coefficients (s_v) for the water column along with date, time, position (latitude, longitude, instrument depth),

temperature, salinity, fluorescence, turbidity, and other sensor data.

Zooplankton net sampling

Two 1 m² MOCNESS (Wiebe et al. 1985) tows were analyzed to identify, enumerate, and measure the zooplankton present in the waters of Wilkinson Basin. A MOCNESS system consists of a series of nine nets, which enables specific depth strata to be sampled. Generally, net 0 was open from the surface to the deepest point of the tow (10–20 m above the bottom), and the remaining nets (1–8) were opened and closed in succession every 25 to 50 m during the return to the surface. The MOCNESS system also recorded the volume of water filtered by each net, the time that each net was opened and closed, depth, salinity, temperature, density, and fluorescence.

The nets were equipped with 333 µm mesh and cod-end buckets for collection of zooplankton and larval fish. Each cod-end sample was split and stored in a buffered formalin solution. Postprocessing of the samples consisted of silhouette photography of the animals (Davis and Wiebe 1985). These images were then examined under a microscope and the organisms were measured and identified by taxonomic group. Numerical density and biomass (mg·m⁻³) were then calculated for each net for each taxonomic group (Davis and Wiebe 1985; Wiebe 1988).

Scattering models

Mathematical models that combine scattering physics and the geometry of the animal shape for several types of zoo-plankton were used to provide backscatter information for single animals for use in the FP analysis (Table 2). These models have been developed previously and only slight modifications to some input parameters have been made in this work. These modifications were limited to body length—width relationships and the use of a simple fluid-like tissue model for certain gelatinous animals. The models represent the three main taxonomic types of zooplankton: fluid-like, elastic-shelled, and gas-bearing animals (Stanton et al. 1998).

Biological scatterers

Fluid-like models were used for copepods, euphausiids, amphipods, and other animals that have a thin shell (which does not support a shear wave) and a body composition that has similar density and sound speed to that of sea water. Fluid-like animals, which constituted the majority of zooplankton taxa that were encountered in the Gulf of Maine and Georges Bank regions, were modeled as weakly scattering, bent, fluid cylinders (Stanton et al. 1993b; Chu et al. 1993). The model has input parameters of animal size (typically, a, the radius), the acoustic frequency (f) or wave number $(k = \frac{2\pi f}{c})$, where c is the speed of sound in seawater), the ratio of sound speed and density between the scatterer and the surrounding fluid (g and h, respectively), the lengthwidth ratio of the animal (β_D) , and the orientation of the cylinder relative to the acoustic wave front. An assumed range of orientation angles based on previous studies was used in modeling the euphausiids, as acoustic backscatter strength is a function of animal orientation (Sameoto 1980; McGehee et al. 1998; Warren et al. 2002). These animals

Scatterer	Scattering model citations	Scattering model parameters
Euphausiids	Stanton et al. 1993 <i>a</i> , Stanton and Chu 2000	$\beta_D = 5.3576$, $R = 0.058$, "head-up" orientation distribution ($20^\circ \pm 20^\circ$)
Copepods and larval crustaceans	Stanton et al. 1993 <i>a</i> , Stanton and Chu 2000	$\beta_D = 2.5497$, $R = 0.058$, uniform orientation distribution (0°–360°)
Amphipods	Stanton et al. 1993 <i>a</i> , Stanton and Chu 2000	$\beta_D = 3.0021$, $R = 0.058$, uniform orientation distribution (0°–360°)
Polychaetes and chaetognaths	Stanton et al. 1993 <i>a</i> , Stanton and Chu 2000	$\beta_D = 17.151$, $R = 0.058$, uniform orientation distribution (0°–360°)
Limacina pteropods	Stanton et al. 1994	R = 0.5
Siphonophore nectophores	Monger et al. 1998	R = 0.028
Siphonophore pneumatophores	Anderson 1950	
Salps	Monger et al. 1998	R = 0.028
Microstructure	Seim 1999	

Table 2. Citations and parameters for the acoustic backscattering models used in the forward problem calculations.

Note: β_D is the length-width ratio (*L/D*) of the animal, *R* is the reflection coefficient.

often orient in a slight head-upward posture and were modeled with a $20^{\circ} \pm 20^{\circ}$ orientation distribution, where 0° is broadside orientation.

Modeling of other fluid-like animals was similar to that for the euphausiids. The value of β_D was changed slightly to better reflect the body shapes of the other fluid-like animals (Table 2). Although Benfield et al. (2001) indicated that copepods may tend to orient themselves vertically in the water column, it was not known under what conditions this occurs, so an average over all orientations (uniform distribution) was used for all fluid-like animals except for euphausiids. Small changes in the values of g and h can cause large variations in the level of scattering from an animal (Chu et al. 2000; Chu and Wiebe 2005), so to minimize variability in this analysis, constant values of g =1.0357 and h = 1.0279 (Foote 1990) were used for all fluidlike animals. It is not known whether the fluid-like animals found in this region have similar material properties as few data are available for animals other than copepods and euphausiids. If differences in the material properties exist for the fluid-like animals, this would cause larger variations in the predicted level of biologically caused backscatter.

Elastic-shelled models were used for animals with hard, elastic shells such as pteropods. Pelagic pteropods are typically very small (<1 mm in diameter) but scatter a large amount of sound (per unit biomass) because of their dense shell. Other strong scatterers are gas-bearing animals such as siphonophores, where the scattering is caused by small gas bubbles used for buoyancy. Gelatinous animals (e.g., salps or medusae) or parts of animals (e.g., siphonophore nectophores) have not been modeled as thoroughly as the fluid-like or elastic-shelled animals. However, because of their body composition, it is believed that they scatter smaller amounts of sound by a mechanism similar to that of the fluid-like animals (Monger et al. 1998).

Several of these zooplankton models have been compared with measured scattering from individual animals (copepods, euphausiids, pteropods, and siphonophore nectophores and pneumatophores) (Stanton and Chu 2000; Stanton et al. 2000; Warren et al. 2001). The remaining zooplankton models have not been tested specifically against measure-

ments from individual animals (amphipods, salps, polychaetes, chaetognaths, larval crustaceans, and cyphanautes); however, these groups contain animals that are believed to be less important acoustically in this study because of either low numerical densities or very weak scattering characteristics.

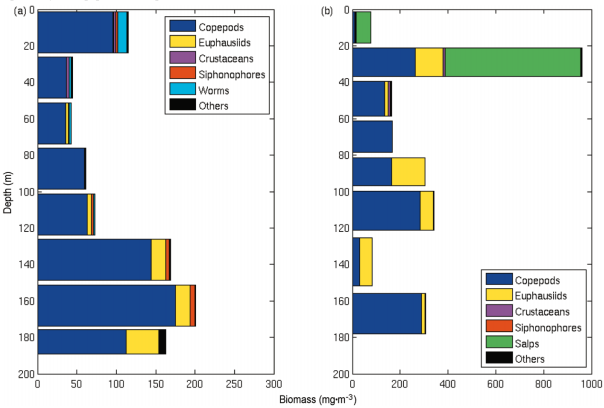
The zooplankton backscatter models were combined with the abundance data from the MOCNESS tows to estimate the amount of biologically caused scattering in the water column. For each animal type collected in a net tow, the backscatter contribution for an individual animal was determined using the appropriate scattering model. These contributions were summed over all animals collected and then divided by the volume of water sampled by the net to arrive at a volume backscatter coefficient for the depth stratum of the net. The contributions from all of the zooplankton were summed and the result was a predicted volume backscatter coefficient for biological sources.

Physical scattering processes

Additional sources of backscatter that have been observed in the vicinity of Georges Bank include suspended sediments and bubbles. These scatterers are not believed to be important in this study because of the absence of sediment in net tows and the relatively calm sea state during the survey period. However, internal waves were seen in the acoustic record during the survey and thus the importance of backscatter from the resultant turbulent microstructure was examined.

Scattering from turbulent microstructure in the water column was analyzed in a parallel manner to that from zoo-plankton except that hydrographic data are used as the scattering model input instead of net tow data. Predictions from the theoretical backscatter model were made based on inputs of temperature, salinity, and the dissipation rates of turbulent kinetic energy and temperature variance (Seim 1999). The latter two values were estimated using temperature and salinity profiles from CTD casts taken either before or after the MOCNESS tow (Table 1) (Thorpe and Brubaker 1983; Warren et al. 2003). Although the ESS system on the MOCNESS provided temperature and salinity profiles, the

Fig. 1. Total biomass estimated from (*a*) MOCNESS 7 and (*b*) MOCNESS 9 data with the relative contributions of the different taxonomic groups. Several taxa are grouped together in the plot, including amphipods and other crustaceans, including larvae ("Crustaceans"), polychaetes and chaetognaths ("Worms"), and pteropods and gelatinous zooplankton ("Others"). The vertical thickness of the bar corresponds to the depth range sampled. For MOCNESS 7, the lower depths were composed primarily of copepods and euphausiids with some gelatinous animals, whereas the surface layers also contained small amounts of polychaetes, chaetognaths, and siphonophore fragments. MOCNESS 9 sampled a large subsurface layer of salps (over 2500 animals·m⁻³), which dominated the biomass sample. The remaining nets were composed primarily of copepods and euphausiids.



sampling rate was limited to 0.25 Hz, thus to resolve temperature and salinity variations at vertical scales less than 1 m, the higher resolution CTD cast data (sampled at 24 Hz) were used.

The CTD data were not collected concurrently with the acoustic and net tow data so there are potential errors in using the CTD data to describe the structure of the water column when the net and acoustic data were recorded. However, hydrographic profiles for each CTD cast were consistent with the profile recorded by the corresponding MOCNESS tow. Although this method is far from ideal for measuring values of turbulent kinetic energy and temperature variance, this method has been used previously to make realistic estimates that compare favorably to measurements of turbulent kinetic energy and temperature variance made in a similar region (Warren et al. 2003; Seim 1999). Sea state was relatively calm during these tows so we believe that errors due to vertical ship and CTD sensor movement are minimal. The estimated level of backscatter from microstructure was then averaged over the depth ranges sampled by each MOCNESS net so that it could be compared with the FP estimates from the zooplankton.

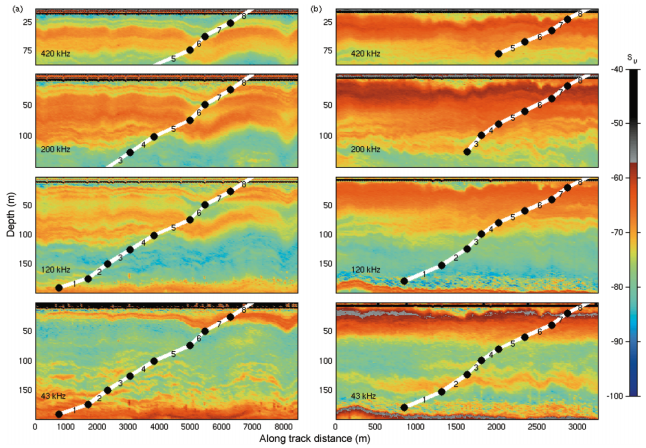
Results

Data are presented for two MOCNESS tows that occurred

in nearby regions but differed in the types of zooplankton present, levels of acoustic backscatter, and water column structure. MOCNESS 7 was lowered to a depth of 191 m and was brought to the surface with a net closed and new net opened at 175, 150, 125, 101, 74, 50, 26, and 0 m. The lower nets contained large amounts of biomass (150–200 mg·m⁻³) and were dominated by copepods and euphausiids (Fig. 1*a*). The surface layer (0–26 m) had a higher level of biomass (over 100 mg·m⁻³) than the other upper water column samples and was composed of copepods, polychaetes, chaetognaths, and amphipods.

Data from MOCNESS 9 show a different depth distribution of biomass, as well as a slightly more diverse taxonomic composition (Fig. 1b). Nets were opened and closed at depths of 180, 153, 124, 99, 80, 60, 39, 20, and 0 m. The zooplankton collected were dominated by an enormous number of salps (2500 animals·m⁻³) near the surface (from depths of 20–40 m), resulting in a large amount of biomass, nearly 1 g·m⁻³. There was also a substantial amount of biomass from 80–124 m that was composed of copepods and euphausiids, as well as a copepod-dominated bottom layer. The salp surface layer was an unusual occurrence on this cruise, and no other net sample from the nine MOCNESS tows collected during the cruise had such a large amount of biomass. For both net tows, the dominant component of biomass at most depths were calanoid copepods.

Fig. 2. BIOMAPER-II echograms for 43, 120, 200, and 420 kHz for (*a*) MOCNESS 7 and (*b*) MOCNESS 9. The white line indicates the path of the net system; black circles indicate where nets were opened and closed. Different regions of the water column have different measured backscattering strengths for the various frequencies. Remnants of an internal wave (undulating backscattering layers) were observed during MOCNESS 7, whereas MOCNESS 9 measured a strong near-surface layer at a depth of 20 m, which is seen most strongly in the 43 kHz echogram.



As each MOCNESS tow was being conducted, BIO-MAPER-II collected acoustic data while being towed at a depth of approximately 5 m beside the ship. The acoustic data were offset horizontally from the MOCNESS samples by the amount of wire out on the net tow (at most a few hundred metres). To compare the acoustic regions with the MOCNESS information, the trajectory of the MOCNESS was overlaid on the acoustic plot to determine where each net sampled (Fig. 2).

The acoustic data for MOCNESS 7 shows strong backscatter at the higher frequencies and weaker backscatter at 43 kHz for much of the water column, although this pattern is reversed for the surface and deepest waters sampled (Fig. 2a). Remnants of an internal wave were observed in the echogram, and the upper layer of the wave, sampled by nets 7 and 8, had the strongest backscatter at the lowest acoustic frequency, whereas backscatter from the thick layer between 50 and 100 m was strongest at the highest frequencies. The echogram collected for MOCNESS 9 showed a midwater scattering layer that was sampled by nets 4-6, with backscatter that had a similar relationship between scattering strength and acoustic frequency (Fig. 2b). This frequency dependence is consistent with the backscatter model used for fluid-like scatterers (Warren et al. 2003). A nearsurface scattering layer (sampled by net 7) shows the opposite effect (strongest backscatter at lower frequencies), which indicates that the scattering was dominated by a different type of scatterer, possibly the large amount of salps or physical processes occurring at the thermocline.

The water column profile for MOCNESS 7 showed a well-mixed region from 20–60 m with a steep temperature and salinity gradient above this layer and a shallower gradient below (Fig. 3a). These mixing processes likely contributed to the backscatter observed between 15 and 100 m in the echogram (Fig. 2a). The hydrographic data for MOCNESS 9 showed a well-mixed region in the upper 20 m of the water column, with a large gradient in temperature, salinity, and density occurring in the next 10 m (Fig. 3b). There were several regions of potential or recent mixing (shown by unstable or nearly vertical sections of the density profile) occurring between 0–20 m, 60–100 m, and 140–180 m, although there were smaller instabilities that occurred throughout the profile.

When FP predictions of backscatter were examined for the individual contributions for different animals or processes, the dominant scatterers for MOCNESS 7 were turbulent microstructure, euphausiids, and siphonophore pneumatophores; however, the amphipod category (which included other larval crustaceans) and chaetognaths also caused appreciable amounts of backscatter (Fig. 4). The

Fig. 3. Temperature (solid line), salinity (broken line), and density (dotted line) profiles collected by the ESS system onboard (*a*) MOCNESS 7 and (*b*) MOCNESS 9. Regions of potential mixing and turbulent microstructure are indicated by nearly vertical or unstable sections of the density profile, e.g., 30–80 m for MOCNESS 7 and 0–20 m, 60–100 m, and 140–180 m for MOCNESS 9.

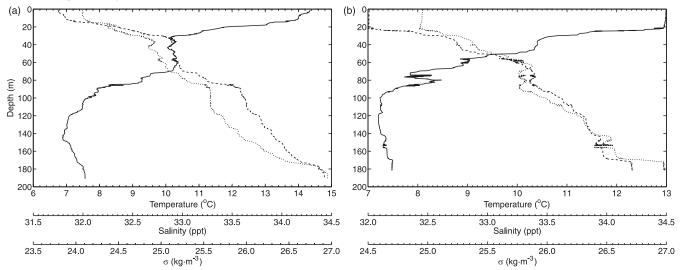


Fig. 4. Forward problem (FP) calculations for each class of scatterer for MOCNESS 7. The different acoustic frequencies (43, 120, 200, and 420 kHz) are represented by squares, stars, circles, and diamonds, respectively. Small copepods had body lengths less than 2.5 mm. Data points that lie above the diagonal line indicate that the FP underestimates the scattering, whereas points below the line are overestimates. Microstructure, siphonophore pneumatophores, chaetognaths, and euphausiids are the strongest contributors to the predicted levels of back-scattering.

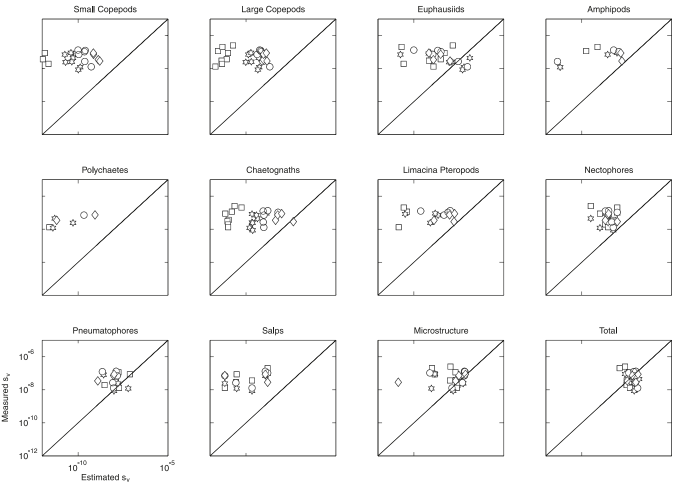
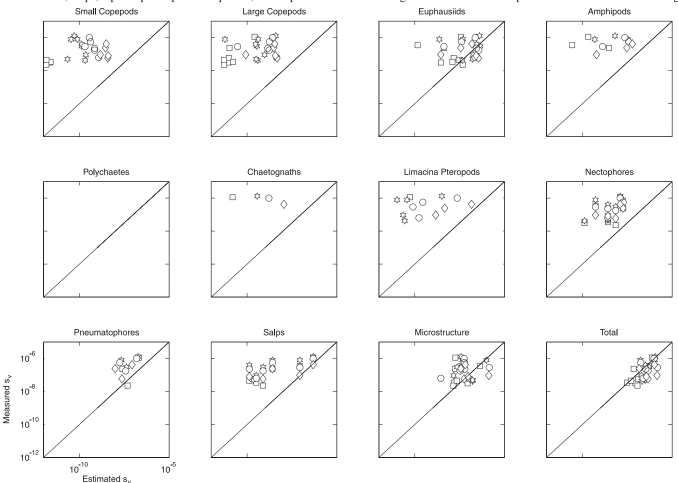


Fig. 5. Forward problem (FP) calculations for each class of scatterer for MOCNESS 9. The different acoustic frequencies (43, 120, 200, and 420 kHz) are represented by squares, stars, circles, and diamonds, respectively. Small copepods had body lengths less than 2.5 mm. Data points that lie above the diagonal line indicate that the FP underestimates the scattering, whereas points below the line are overestimates. Microstructure, salps, siphonophore pneumatophores, and euphausiids are the strongest contributors to the predicted levels of backscattering.



other animals (particularly the abundant copepods) contributed little to the overall predicted scattering except at the highest frequencies. Copepods also contributed little to the predictions for MOCNESS 9. Turbulence, euphausiids, salps, and siphonophore pneumatophores were the largest contributors to the backscatter (Fig. 5). It is striking that the copepods that were by far the largest contributors to biomass have such a small contribution to the predicted levels of backscatter. This is primarily a function of copepod size (a few millimetres in length) and acoustic frequency or wavelength. For the frequencies used in this survey, copepod backscatter is primarily a function of animal size, and despite their numerical abundance in the net tow data, they are simply too small to contribute much to the predicted level of backscatter except at the higher frequencies.

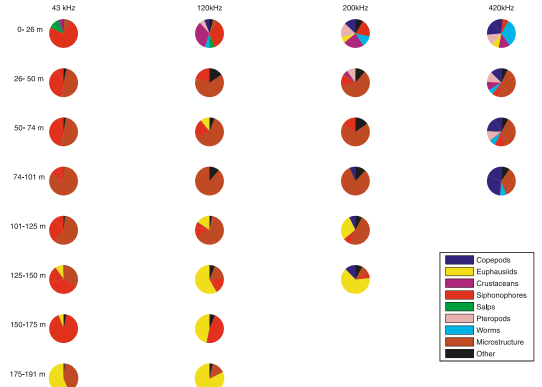
The MOCNESS data provided information about the contributions that different zooplankton taxa make to the overall amount of biomass. Similarly, the relative contributions of different biological and physical sources to the total amount of predicted backscatter in the water column can be made by combining MOCNESS data, CTD data, and backscatter models. The relative contribution of each scattering source (each animal taxa and microstructure) was calculated for

each MOCNESS net depth range and BIOMAPER-II frequency (Figs. 6 and 7). The percentage contribution to the total predicted backscatter strength was found by dividing the predicted volume backscatter coefficient for each scatterer type by the overall calculated backscatter prediction. The percentage of total predicted scattering from physical (nonbiological) sources was calculated. The measured level of scattering (from the BIOMAPER-II data) was then reduced by this percentage to arrive at a corrected amount of measured backscatter that is believed to be from biological scatterers.

For example, the measured level of backscatter for MOCNESS 9 from depths of 0–20 m at 120 kHz is $s_{\rm v}$ = 1.58 × 10⁻⁶ m⁻¹ ($S_{\rm v}$ = –58.0 dB). From Fig. 7, only 8% of the predicted backscatter for this sample is from biological sources. By multiplying the percentage of biologically caused predicted backscatter and the measured level of backscatter, an estimate of the biologically caused scatter in the water column was made, $s_{\rm v}$ = 1.26 × 10⁻⁷ m⁻¹ ($S_{\rm v}$ = –69.0 dB).

Nonbiological backscatter contributions were important for several regions sampled by MOCNESS 7 (Fig. 6). The predicted contributions from microstructure were largest for

Fig. 6. Percentage breakdown of forward problem calculations for each net and acoustic frequency of MOCNESS 7. Turbulent microstructure contributes large amounts to the total backscattering in the midwater regions, whereas euphausiids backscatter a majority of the sound in the deeper water. Siphonophores contribute greatly at the lower frequencies, but not at the higher ones. The surface region (0–20 m) has a very diverse group of scatterers.



the region between 20 and 100 m (which again corresponded to regions of mixing indicated in the hydrographic data), whereas euphausiids were the main scatterers for the deepest nets. Siphonophores contributed to the backscatter more for lower acoustic frequencies and were negligible at the highest frequency. The backscatter in the near surface was the most diverse with regard to scatterer type, with nearly all taxonomic types contributing.

For MOCNESS 9, the deeper water column and surface layer were dominated by scattering from microstructure (Fig. 7). These large backscatter contributions from turbulent microstructure occurred in the same regions that the hydrographic profile data indicated was well mixed (0–20 m and 140–170 m). Euphausiids were the dominant scatterers in the midwater depths, with siphonophore pneumatophores and nectophores also contributing. Salps were extremely weak scatterers, and although outnumbering the other animals and dominating the biomass in the near-surface region, they contributed only 30%–60% to the total backscatter in that region.

If the FP is well posed and one taxa dominates the measured scattering, then there should be a relationship between biomass and measurements of backscatter strength. Both biomass and volume backscatter cross sections (s_v) are linear functions (for a particular taxa) of the number of animals. Therefore it is likely that a relationship between biomass and volume scattering strength should exist. The relationship between biomass and measured backscatter may not be linear, however, if more than one scattering process (or

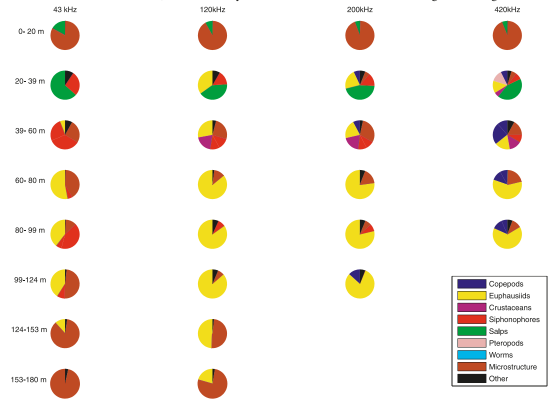
taxonomic type or size class) is substantially contributing to the measured backscatter.

The biomass and acoustic backscatter data sets from both MOCNESS tows were combined, and the regression between the logarithm of biomass and measured acoustic backscatter strength (S_v , a logarithmic measure of acoustic backscatter) was found for each acoustic frequency. The log of both biomass and backscatter was used as some of the acoustic data (specifically predicted backscatter for some scatterer types) ranged over nearly five orders of magnitude. The backscatter model used for salps has not been as well tested, by comparing theoretical backscatter predictions with measured backscatter from individual animals, as the backscatter models used for other animals. Because of this fact and the extremely high biomass of salps caught in net 7 of MOCNESS 9, the data from this net were not included in this analysis.

There was not a strong relationship (all r^2 values < 0.4) between log-transformed biomass and measured backscatter levels for any of the four frequencies used (Fig. 8). A poor relationship between zooplankton biomass and acoustic backscatter is likely to occur when nonzooplankton scatterers are contributing to the measured amount of acoustic backscatter. This result was not surprising as turbulent microstructure was predicted to contribute greatly to the measured backscatter for some portions of the water column.

A similar analysis was performed for biomass and biologically caused backscatter (Fig. 9). When backscatter attributed to physical processes was removed, the relation-

Fig. 7. Percentage breakdown of forward problem calculations for each net and acoustic frequency of MOCNESS 9. Euphausiids dominate the backscattering in the midwater depths, whereas turbulence contributes strongly near both the bottom and the surface. The salps, which dominated the biomass in the near-surface net, contribute only 30% to 60% to the total backscattering for that region.



ship between log measures of biomass and acoustic back-scatter was more linear. Regression coefficients improved for all frequencies, indicating a better correlation between biomass and backscatter. It must be noted that the regression coefficients for each frequency are still fairly small (r^2 ranged from 0.38 to 0.52); however, these values are a factor of two or three larger than if the source of the backscatter is not identified. By accounting for the source of acoustic backscatter using hydrographic and net tow information, this method can be used to improve the use of acoustic backscatter data as a measure of zooplankton biomass.

Discussion

One of the goals of acoustic surveys is to estimate zooplankton biomass, and this requires that the relationship between biomass and acoustic backscatter is well understood. The data presented here indicate that improvements can be made in the interpretation of field-collected survey data if the contributions of all scattering sources are quantified. The relative importance of physical and biological sources of acoustic backscatter will vary with location in the ocean, and certainly, some regions will not have substantial backscatter from physical processes in the water column, whereas other areas (such as the sites in this study that have internal waves present) will have significant contributions to the backscatter from nonbiological sources. The modeling efforts outlined in this work provide one way of determining if physical sources of backscatter will need to be accounted for when interpreting acoustic backscatter survey data. However, these improvements are just one step of many that need to be taken in order that acoustic surveys may provide estimates of zooplankton abundance that are accurate and ecologically useful. Given that biomass and predictions of biologically caused backscatter are not perfectly correlated, sources of error in the analysis, such as inaccuracies in the backscatter models used, must be examined.

The zooplankton backscatter models for fluid-like, elastic-shelled, and gas-bearing zooplankton have been used previously in the analysis of field-collected data (Wiebe et al. 1996, 1997; Greene et al. 1998); however, there are many variables used in these models that are inadequately understood, such as animal behavior and orientation or the material properties (g and h) of the zooplankton. A better understanding of the scattering model inputs would reduce errors associated with these types of animals. Furthermore, there are numerous types of animals (salps, polychaetes, chaetognaths, gelatinous zooplankton) with backscatter characteristics that have been neither modeled nor measured in a laboratory environment.

Uncertainty about the inputs to the backscatter models is a concern for the microstructure model as well. Proper instrumentation was not present to measure the dissipation rates of turbulent kinetic energy and temperature variance, which are vital inputs into the theoretical microstructure of backscatter models, so the method used to estimate these inputs was not ideal. Although this method provided reasonable estimates of ϵ and χ , it likely overestimated the scattering contributions from microstructure. For example, some

Fig. 8. The relationship between the logarithm of biomass and measurements of acoustic backscatter strength for MOCNESS 7 (squares) and MOCNESS 9 (circles), excluding net 7 from MOCNESS 9. Most frequencies show little correlation between these two variables, except for the highest frequency (420 kHz).

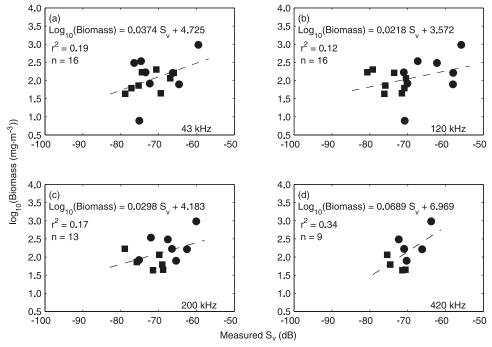
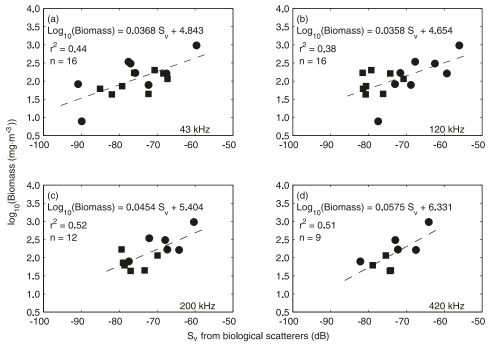


Fig. 9. The relationship between the logarithm of biomass and predictions of biologically caused acoustic backscatter for MOCNESS 7 (squares) and MOCNESS 9 (circles), excluding net 7 from MOCNESS 9. All of the acoustic frequencies show improved regressions and fairly linear relationships for the data.



regions of MOCNESS 9 have microstructure-caused s_v values that were larger than the backscatter measured by BIOMAPER-II. Further complicating this issue is the possibility that the vertical migration of animals may be creating significant amounts of turbulence and mixing (Huntley and Zhou 2004; Kunze et al. 2006).

Other possible sources of error in the FP analysis include erroneous zooplankton abundance and composition data and inaccurate measurements of the acoustic backscatter. Net tow information from MOCNESS systems has been used for several decades, and sampling errors from it are likely limited to net avoidance by large zooplankton (Wiebe et al.

2004) and gelatinous animals being destroyed by the net mesh. Finally, the undersampling of animals by either nets (e.g., large euphausiids or small fish) or lower acoustic frequencies (e.g., copepods) will cause errors in the FP analysis. One approach that has been used in the analysis of acoustic scattering data (Warren et al. 2003) is to use the net tow estimates of numerical density as a lower bound on the true value (as you cannot have more animals in a net than are present in the water column) and to use the acoustically inferred estimates of biomass as an upper bound (as these rely on measures of backscatter strength that likely contain contributions from other scattering sources). In this manner, combining acoustic and net tow data can provide an upper and lower estimate of the abundance of zooplankton in the water column.

This study also demonstrates the importance of resolving the changes in biological and physical backscatter sources within the water column. A multiple net system, or other methods such as video or optical ground-truthing, may be necessary to accurately assess acoustic surveys of zooplankton biomass, particularly where the taxonomic components of the zooplankton community are diverse. Providing this vertical resolution and ground-truthing of the acoustic data also allows us to observe partitioning of the water column into different habitats that would not be apparent from either the acoustic data alone or a vertically integrating net tow. Although some regions of the ocean do have patches with a single dominant taxa (e.g., Antarctic krill), the variation in abundance and distribution of zooplankton taxa observed over a 200 m vertical span in this study demonstrates the importance of measuring and quantifying these changes.

The difference between predictions of $S_{\rm v}$ and those values that would perfectly correlate with the biomass data are on the order of 5–10 dB (assuming that the biomass data are accurate). These differences become very large when back-scatter strengths are converted to estimates of biomass; therefore, these predictions result in estimates of zooplank-ton biomass that are correct to roughly an order of magnitude. In certain cases, this level of error may be acceptable, but further work is needed to reduce this uncertainty. Without accounting for the source of acoustic scattering in the water column, estimates of biomass from acoustics are likely to have even larger errors.

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References

Anderson, V.C. 1950. Sound scattering from a fluid sphere. J. Acoust. Soc. Am. 22: 426–431. doi:10.1121/1.1906621.Benfield, M.C., Davis, C.S., and Gallager, S.M. 2001. Estimating

- the in-situ orientation of *Calanus finmarchicus* on Georges Bank using the Video Plankton Recorder. Plankton Biol. Ecol. **47**: 69–72.
- Benfield, M.C., Lavery, A.C., Wiebe, P.H., Greene, C.H., Stanton, T.K., and Copley, N.J. 2003. Distributions of physonect siphonulae in the Gulf of Maine and their potential as important sources of acoustic scattering. Can. J. Fish. Aquat. Sci. 60: 759–772. doi:10.1139/f03-065.
- Brierley, A.S., Ward, P., Watkins, J.L., and Goss, C. 1998. Acoustic discrimination of Southern Ocean zooplankton. Deep-Sea Res. Part II Top. Stud. Oceanogr. 45: 1155–1173. doi:10.1016/ S0967-0645(98)00025-3.
- Chu, D., and Wiebe, P.H. 2005. Measurements of sound speed and density contrasts of zooplankton in Antarctic waters. ICES J. Mar. Sci. 62: 818–831. doi:10.1016/j.icesjms.2004.12.020.
- Chu, D., Foote, K.G., and Stanton, T.K. 1993. Further analysis of target strength measurements of Antarctic krill at 38 and 120 kHz: comparison with deformed cylinder model and inference of orientation distribution. J. Acoust. Soc. Am. 93: 2985–2988. doi:10.1121/1.405818.
- Chu, D., Wiebe, P.H., and Copley, N. 2000. Inference of material properties of zooplankton from acoustic and resistivity measurements. ICES J. Mar. Sci. 57: 1128–1142. doi:10.1006/jmsc. 2000.0800.
- Davis, C.S., and Wiebe, P.H. 1985. Macrozooplankton biomass in a warm-core Gulf Stream ring: time series changes in size, structure, and taxonomic composition and vertical distribution. J. Geophys. Res. 90: 8871–8884. doi:10.1029/JC090iC05p08871.
- Davis, C.S., Gallager, S.M., Marra, M., and Stewart, W.K. 1996. Rapid visualization of plankton abundance and taxonomic composition using the Video Plankton Recorder. Deep-Sea Res. 43: 1947–1970. doi:10.1016/S0967-0645(96)00051-3.
- Foote, K.G. 1990. Speed of sound in *Euphausia superba*. J. Acoust. Soc. Am. **87**: 1405–1408. doi:10.1121/1.399436.
- Foote, K.G., and Stanton, T.K. 2000. Acoustical methods. *In ICES* zooplankton methodology manual. *Edited by R. Harris*, P.H. Wiebe, J. Lenz, H.R. Skjoldal, and M. Huntley. Academic Press, London. pp. 223–258.
- Goodman, L. 1990. Acoustic scattering from ocean microstructure. J. Geophys. Res. 95: 11557–11573. doi:10.1029/JC095iC07p11557.
- Greene, C.H., Wiebe, P.H., Pershing, A.J., Gal, G., Popp, J.M., Copley, N.J., Austin, T.C., Bradley, A.M., Goldsborough, R.G., Dawson, J., Hendershott, R., and Kaartvedt, S. 1998. Assessing the distribution and abundance of zooplankton: a comparison of acoustic and net-sampling methods with D-BAD MOCNESS. Deep-Sea Res. Part II Top. Stud. Oceanogr. 45: 1219–1237. doi:10.1016/S0967-0645(98)00033-2.
- Hewitt, R.P., and Demer, D.A. 2000. The use of acoustic sampling to estimate the dispersion and abundance of euphausiids, with an emphasis on Antarctic krill, *Euphausia superba*. Fish. Res. **47**: 215–229. doi:10.1016/S0165-7836(00)00171-5.
- Holliday, D.V., and Pieper, R.E. 1995. Bioacoustical oceanography at high frequencies. ICES J. Mar. Sci. 52: 279–296. doi:10.1016/ 1054-3139(95)80044-1.
- Huntley, M.E., and Zhou, M. 2004. Influence of animals on turbulence in the sea. Mar. Ecol. Prog. Ser. 273: 65–79. doi:10.3354/meps273065.
- Kunze, E., Dower, J.F., Beveridge, I., Dewey, R., and Bartlett, K.P. 2006. Observations of biologically generated turbulence in a coastal inlet. Science (Washington, D.C.), 313: 1768–1770. doi:10.1126/science.1129378. PMID:16990545.
- Lavery, A.C., Schmitt, R.W., and Stanton, T.K. 2003. High-frequency acoustic scattering from turbulent oceanic microstructure: the importance of density fluctuations. J. Acoust.

Soc. Am. **114**: 2685–2697. doi:10.1121/1.1614258. PMID: 14650005.

- Mair, A.M., Fernandes, P.G., Lebourges-Dhaussy, A., and Brierley, A.S. 2005. An investigation into the zooplankton composition of a prominent 38-kHz scattering layer in the North Sea. J. Plankton Res. 27: 623–633. doi:10.1093/plankt/fbi035.
- Martin, L.V., Stanton, T.K., Wiebe, P.H., and Lynch, J.F. 1996. Acoustic classification of zooplankton. ICES J. Mar. Sci. 53: 217–224. doi:10.1006/jmsc.1996.0025.
- McGehee, D.E., O'Driscoll, R.L., and Martin-Traykovski, L.V. 1998. Effects of orientation on acoustic scattering from Antarctic krill at 120 kHz. Deep-Sea Res. Part II Top. Stud. Oceanogr. 45: 1273–1294. doi:10.1016/S0967-0645(98)00036-8.
- Monger, B.C., Chinniah-Chandy, S., Meir, E., Billings, S., Greene, C.H., and Wiebe, P.H. 1998. Sound scattering by the gelatinous zooplankters *Aequorea victoria* and *Pleurobrachia bachei*. Deep-Sea Res. Part II Top. Stud. Oceanogr. **45**: 1255–1271. doi:10.1016/S0967-0645(98)00029-0.
- Orr, M.H., Haury, L.R., Wiebe, P.H., and Briscoe, M.G. 2000. Backscatter of high-frequency (200 kHz) acoustic wavefields from ocean turbulence. J. Acoust. Soc. Am. **108**: 1595–1601. doi:10.1121/1.1286883. PMID:11051487.
- Ross, T., and Lueck, R. 2005. Estimating turbulent dissipation rates from acoustic backscatter. Deep-Sea Res. 52: 2353–2365. doi:10.1016/j.dsr.2005.07.002.
- Sameoto, D.D. 1980. Quantitative measurements of Euphausiids using a 120-kHz sounder and their in situ orientation. Can. J. Fish. Aquat. Sci. 37: 693–702. doi:10.1139/f80-087.
- Seim, H.E. 1999. Acoustic backscatter from salinity microstructure. J. Atmos. Ocean. Tech. 16: 1491–1498. doi:10.1175/1520-0426(1999)016<1491:ABFSM>2.0.CO;2.
- Seim, H.E., Gregg, M.C., and Miyamoto, R.T. 1995. Acoustic back-scatter from turbulent microstructure. J. Atmos. Ocean. Tech. 12: 367–380. doi:10.1175/1520-0426(1995)012<0367:ABFTM>2.0. CO;2.
- Stanton, T.K., and Chu, D. 2000. Review and recommendations for modeling of acoustic scattering by fluid-like elongated zooplankton: euphausiids and copepods. ICES J. Mar. Sci. 57: 793– 807. doi:10.1006/jmsc.1999.0517.
- Stanton, T.K., Chu, D., Wiebe, P.H., and Clay, C.S. 1993a. Average echoes from randomly oriented random-length finite cylinders: zooplankton models. J. Acoust. Soc. Am. 94: 3463– 3472. doi:10.1121/1.407200.
- Stanton, T.K., Clay, C.S., and Chu, D. 1993b. Ray representation of sound scattering by weakly scattering deformed fluid cylinders: simple physics and application to zooplankton. J. Acoust. Soc. Am. **94**: 3454–3462. doi:10.1121/1.407199.
- Stanton, T.K., Wiebe, P.H., Chu, D., Benfield, M.C., Scanlon, L., Martin, L., and Eastwood, R.L. 1994. On acoustic estimates of zooplankton biomass. ICES J. Mar. Sci. 51: 505–512. doi:10. 1006/jmsc.1994.1051.
- Stanton, T.K., Chu, D., and Wiebe, P.H. 1998. Sound scattering by several zooplankton groups. II. Scattering models. J. Acoust. Soc. Am. 103: 236–253. doi:10.1121/1.421110. PMID:9440326.

- Stanton, T.K., Chu, D., Wiebe, P.H., Eastwood, R.L., and Warren, J.D. 2000. Acoustic scattering by benthic and pelagic shelled animals. J. Acoust. Soc. Am. 108: 535–550. doi:10.1121/1. 429584. PMID:10955618.
- Thorpe, S.A., and Brubaker, J.M. 1983. Observations of sound reflection by temperature microstructure. Limnol. Oceanogr. 28: 601–613.
- Trevorrow, M.V. 1998. Observations of internal solitary waves near the Oregon coast with an inverted echo sounder. J. Geophys. Res. **103**: 7671–7680. doi:10.1029/98JC00101.
- Warren, J.D., Stanton, T.K., Benfield, M.C., Wiebe, P.H., Chu, D., and Sutor, M. 2001. In situ measurements of acoustic target strengths of gas-bearing siphonophores. ICES J. Mar. Sci. 58: 740–749. doi:10.1006/jmsc.2001.1047.
- Warren, J.D., Stanton, T.K., McGehee, D.E., and Chu, D. 2002. Effect of animal orientation on acoustic estimates of zooplankton properties. IEEE J. Oceanic Eng. 27: 130–138. doi:10.1109/ 48.989899.
- Warren, J.D., Stanton, T.K., Wiebe, P.H., and Seim, H.E. 2003. Inference of biological and physical parameters in an internal wave using multiple frequency acoustic scattering data. ICES J. Mar. Sci. 60: 1033–1046. doi:10.1016/S1054-3139(03)00121-8.
- Wiebe, P.H. 1988. Functional regression equations for zooplankton displacement volume, wet weight, dry weight, and carbon: a correction. Fish. Bull. (Washington, D.C.), 86: 833–835.
- Wiebe, P.H., Morton, A.W., Bradley, A.M., Backus, R.H., Craddock, J.E., Barber, V., Cowles, T.J., and Flierl, G.R. 1985. New developments in the MOCNESS, an apparatus for sampling zooplankton and micronekton. Mar. Biol. (Berl.), 87: 313–323. doi:10. 1007/BF00397811.
- Wiebe, P.H., Mountain, D.G., Stanton, T.K., Greene, C.H., Lough, G., Kaartvedt, S., Dawson, J., and Copley, N. 1996. Acoustical study of the spatial distribution of plankton on Georges Bank and the relationship between volume backscattering strength and the taxonomic composition of the plankton. Deep-Sea Res. Part II Top. Stud. Oceanogr. 43: 1971–2001. doi:10.1016/S0967-0645(96)00039-2.
- Wiebe, P.H., Stanton, T.K., Benfield, M.C., Mountain, D.G., and Greene, C.H. 1997. High-frequency acoustic volume backscattering in the Georges Bank coastal region and its interpretation using scattering models. IEEE J. Oceanic Eng. 22: 445–464. doi:10.1109/48.611135.
- Wiebe, P.H., Stanton, T.K., Greene, C.H., Benfield, M.C., Sosik, H.M., Austin, T., Warren, J.D., and Hammer, T. 2002. BIOMA-PER II: an integrated instrument platform for coupled biological and physical measurements in coastal and oceanic regimes. IEEE J. Oceanic Eng. 27: 700–716. doi:10.1109/JOE.2002. 1040951.
- Wiebe, P.H., Ashjian, C., Gallager, S., Davis, C., Lawson, G., and Copley, N. 2004. Using a high powered strobe light to increase the catch of Antarctic krill. Mar. Biol. (Berl.), 144: 493–502. doi:10.1007/s00227-003-1228-z.