Evidence for the distortion product frequency place as a source of distortion product otoacoustic emission (DPOAE) fine structure in humans. I. Fine structure and higher-order DPOAE as a function of the frequency ratio $f_2/f_1$\textsuperscript{a)}

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Critical experiments were performed in order to validate the two-source hypothesis of distortion product otoacoustic emissions (DPOAE) generation. Measurements of the spectral fine structure of DPOAE in response to stimulation with two sinusoids have been performed with normal-hearing subjects. The dependence of fine-structure patterns on the frequency ratio $f_2/f_1$ was investigated by changing $f_1$ or $f_2$ only (fixed $f_2$ or fixed $f_1$ paradigm, respectively), and by changing both primaries at a fixed ratio and looking at different order DPOAE. When $f_2/f_1$ is varied in the fixed ratio paradigm, the patterns of $2f_1-f_2$ fine structure vary considerably more if plotted as a function of $f_2$ than as a function of $f_{DP}$. Different order distortion products located at the same characteristic place on the basilar membrane (BM) show similar patterns for both, the fixed-$f_2$ and $f_{DP}$ paradigms. Fluctuations in DPOAE level up to 20 dB can be observed. In contrast, the results from a fixed-$f_{DP}$ paradigm do not show any fine structure but only an overall dependence of DP level on the frequency ratio, with a maximum for $2f_1-f_2$ at $f_2/f_1$ close to 1.2. Similar stimulus configurations used in the experiments have also been used for computer simulations of DPOAE in a nonlinear and active model of the cochlea. Experimental results and model simulations give strong evidence for a two-source model of DPOAE generation: The first source is the initial nonlinear interaction of the primaries close to the $f_2$ place. The second source is caused by coherent reflection from a re-emission site at the characteristic place of the distortion product frequency. The spectral fine structure of DPOAE observed in the ear canal reflects the interaction of both these sources.

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INTRODUCTION

Narrow-band distortion product otoacoustic emissions (DPOAE) are low-level sinusoids recordable in the occluded ear canal at certain combination frequencies during continuous stimulation with two tones. They are the result of the nonlinear interaction of the tones in the cochlea. In human subjects, DPOAE typically exhibit a pronounced spectral fine structure when varying the frequencies of both primaries simultaneously ($f_1,f_2$) at a fixed frequency ratio $f_2/f_1$ (Gaskill and Brown, 1990; He and Schmiedt, 1993). The variations of DPOAE level with frequency show a periodicity of about 3/32 octaves (He and Schmiedt, 1993; Mauermann et al., 1997b) in a depth up to 20 dB. DPOAE can be recorded in almost any normal-hearing subject and in subjects with hearing loss up to 50 dB HL (Smurzynski et al., 1990). Because of the narrow-band nature of both stimuli and emissions, they provide a frequency-specific method to explore cochlear mechanics. Therefore, DPOAE are of great interest not only in laboratory studies but also as a diagnostic tool for clinical audiology. However, since it is not as yet completely understood which sources along the cochlear partition contribute to the emission measured in the ear canal, the applicability of DPOAE for, e.g., “objective” audiometry is limited at present.

For DPOAE with frequencies below the primary frequencies ($2f_1-f_2,3f_1-2f_2$, etc.), it is widely accepted that the generation site is the overlap region of the excitation patterns of the two primaries, which has a maximum close to the characteristic place around $f_2$. Although the generation of distortion products due to the interaction of the two primaries is in principle spread over the whole cochlea, a region of about 1 mm around the characteristic place of $f_2$ has been suggested to give the maximum contribution (van Hengel and Duifhuis, 1999). This region of maximum contribution is referred to as the $f_2$ site.
It is still a point of discussion whether the generation site is the only source or to what extent other sources might also contribute to the emission. The DPOAE fine structure found in human subjects is closely related to this question of DPOAE sources. The fine structure might reflect local BM properties of either the generation site or of the re-emission site. It could also result from the interference between two or more sources or even from a combination of both local properties and interference effects. Studying the properties of DPOAE fine structure may result in further insight into BM mechanisms and the location of DP sources.

The patterns of DPOAE fine structure get shifted along the frequency axis when the primary levels are increased (He and Schmiedt, 1993; Mauermann et al., 1997b) or the frequency ratio of the primaries is changed (Mauermann et al., 1997a). On one hand, these shifts may cause some problems in the interpretation of DPOAE measurements, especially DPOAE growth functions. Peaks may change to notches or vice versa. This is most probably the reason for the notches found in human DPOAE growth functions (He and Schmiedt, 1993) and is critical for a direct correlation of DPOAE level to hearing threshold. On the other hand, the correct interpretation of these fine-structure shifts can aid in a detailed understanding of BM mechanisms.

He and Schmiedt (1993) showed that the level-dependent shift of DPOAE fine structure is consistent with the results of Ruggero and Rich (1991) on the shift of the maximum basilar-membrane response in the chinchilla. They interpreted the fine structure as an effect of local BM properties in the region of the primaries and the level-dependent shift as a result of the shift of the primary excitation on the BM (He and Schmiedt, 1993; Sun et al., 1994a, b). Varying only one primary level while holding the other fixed causes pattern shifts in different directions, dependent on whether the primary level at \( f_1 \) or at \( f_2 \) is varied. He and Schmiedt (1997) argued that these effects strongly support the idea that the DPOAE fine structure might reflect mechanical properties of the overlapping area of the primary excitation.

However, Heitmann et al. (1998) showed that the fine structure disappears when the DPOAE is measured with a third tone close to the distortion product frequency \( f_{DP} \) as a suppressor. This result is interpreted as evidence for an additional second source around the characteristic place of \( 2f_1-f_2 \), which has a major influence on the fine-structure pattern. The contribution of a second source at the place of \( 2f_1-f_2 \) is also supported by several experiments on DPOAE suppression. Kummer et al. (1995) reported that in some cases a suppressor close to \( 2f_1-f_2 \) results in more suppression than one close to \( f_2 \). Gaskell and Brown (1996) also found that the DP level is still sensitive to a suppressor near \( 2f_1-f_2 \), although the major suppression effects they observed were for suppressor frequencies close to \( f_2 \). Brown et al. (1996) showed “that it may be legitimate to analyze DP as vector sum of two gross components,” (Brown et al., 1996, p. 3263).

Throughout the present paper, experimental results from normal-hearing subjects and computer simulations will be presented examining in detail the properties of DPOAE fine structure for equal-level primaries and varied ratios of the primary frequencies. Three “critical” experiments have been performed, using different experimental paradigms, that aim to clarify where and how DPOAE fine structure is generated. Experiment 1 investigates DPOAE fine-structure patterns for different \( f_2/f_1 \) to determine whether the fine structure is dominated by local properties of the \( f_2 \) region or if a supposed re-emission site around \( f_{DP} \) is of some importance. Experiment 2 is designed to test the influence of the relative phase between the suggested emission sites by investigating the patterns of different order DPOAE. Finally, in experiment 3 the DPOAE patterns from a \( f_{DP} \)-fixed and a \( f_2 \)-fixed paradigm are compared to find out if DPOAE fine structure is mainly influenced by one of these two sites.

In addition to the recordings from subjects, all experimental paradigms were also assessed with a computer simulation of DPOAE using a nonlinear and active transmission line model of the cochlea. This model includes an impedance function as suggested by Zweig (1991) which produces excitation patterns with a broad and tall peak. Within the model, statistical fluctuations of stiffness along the cochlear partition are sufficient to create quasiperiodic OAE fine-structure patterns, as reported by Zweig and Shera (1995).

I. METHODS

A. Subjects

Seven normal-hearing subjects, ranging in age from 25 to 30 years, participated in this study. Their hearing thresholds were better than 15 dB HL for all audiometric frequencies in the range 250 Hz–8 kHz, and none of the subjects had a history of any hearing problems. Spontaneous otoacoustic emissions (SOAE) were observed in only one of the subjects (subject SE). DPOAE were recorded from one ear of each subject during several sessions lasting 60–90 min. During the sessions, the subjects were seated comfortably in a sound-insulated booth (IAC-1200 CT).

B. Instrumentation and signal processing

An insert ear probe, type ER-10C, was used to record DPOAE. The microphone output was connected to a low-noise amplifier, type SR560, and then converted to digital form using the 16-bit A/D converters on a signal-processing board (Ariel DSP-32C) in a personal computer. All stimuli were generated digitally. After D/A conversion by the 16-bit D/A converters on the Ariel board and low-pass filtering (Kemo VBF44, 8.5 kHz) they were presented to the subjects via a computer-controlled audiometer. The DSP was used for on-line analysis and signal conditioning of the recorded emissions, including artifact rejection, averaging in the time domain to improve the signal-to-noise ratio, and fast Fourier transform (FFT). For each signal configuration, at least 16 but usually 256 frames were averaged, using a frame length of 186 ms (4096 samples). If required, the number of averages could be increased during the recording to get a sufficient signal-to-noise ratio for the frequencies of interest.

Two sinusoids were generated as even harmonics of the frame rate (5.38 Hz) at a sampling rate of 22 050 Hz. The tones were presented continuously to the subject. To compensate for the ear-canal transfer function, an individual ad-
justment of the primaries to the desired sound-pressure level of 60 dB SPL was performed automatically before each run, taking into account the transfer function of the probe microphone. The variations of attenuation within and between subjects were approximately within a range of 5 dB.

C. Experimental paradigms

1. Dependence of the $2f_1-f_2$ DPOAE on frequency ratio

In experiment 1, the effect of the frequency ratio $f_2/f_1$ on the fine-structure patterns of the DPOAE at $2f_1-f_2$ was investigated. Fine-structure patterns for this distortion product were recorded in all subjects for seven different frequency ratios $f_2/f_1$, fixed at 1.07, 1.1, 1.13, 1.16, 1.19, 1.22, and 1.25. Due to the additional requirement to select the primaries as harmonics of the frame rate, minor deviations upper to 0.002 from the desired frequency ratio were present. Recordings were taken covering a frequency range of two octaves ($f_1=1-4$ KHz), divided into four sessions covering half an octave each for all of the different frequency ratios specified above. The frequency step between adjacent single recordings was 1/48 octave.

It is assumed that the small changes in the frequency ratio cause only small changes in the fine-structure patterns, i.e., the patterns remain comparable. Consequently, if the fine-structure patterns are mostly influenced by the local properties of the generation site near the characteristic place of $f_2$, the patterns for different $f_2/f_1$ should show a high stability when plotted as a function of $f_2$. If, however, the local properties of a presumed re-emission site near the characteristic place of the DPOAE frequency play a major role, the stability of the patterns should be greater when plotted as a function of $f_{DP}$.

2. Different order DPOAE

In experiment 2, fine-structure patterns for different order DPOAE (e.g., $2f_1-f_2$, $3f_1-2f_2$, and $4f_1-3f_2$) were recorded from six of our seven subjects. The frequency ratios were chosen to get identical frequency pairs either for $f_2$ and $2f_1-f_2$, $f_2$ and $3f_1-2f_2$, and $f_2$ and $4f_1-3f_2$, or for $f_1$ and $2f_1-f_2$, $f_1$ and $3f_1-2f_2$, and $f_1$ and $4f_1-3f_2$, respectively. The condition for identical $f_2$ and $f_{DP}$ frequencies is fulfilled for the frequency ratios $f_2/f_1=1.22$, 1.137, and 1.099. Similarly, the required identity of $f_1$ and $f_{DP}$ frequencies is fulfilled at frequency ratios $f_2/f_1=1.22$, 1.11, and 1.073 (see Fig. 1).

If the DPOAE fine structure is mainly caused by two sources, one at the generation site and the other one at the distortion frequency site, measurements for different order DPOAE with identical $f_2$ and $f_{DP}$ frequencies should result in very similar patterns, since both $f_2$ and the observed DP frequency are the same, i.e., the characteristic places of the two assumed sources and hence the phase relation between the two is almost identical (see Fig. 1, left column). With identical $f_1$ and $f_{DP}$ frequencies, a small variation in the pattern is expected, indicating the influence of the change in the relative phase of the $f_2$ and $f_{DP}$ components (see Fig. 1, right column).

D. Analysis

For further analysis, the frequencies in all experiments were transformed to their characteristic places $x(f)$ on the BM using the place–frequency map proposed by Greenwood (1991). Although the frequencies used are almost equally spaced on the Greenwood map, there are some deviations from this. These deviations are mainly due to the fact that the primaries were selected as harmonics of the frame rate. To compensate for that, the data were interpolated using a cubic spline algorithm (MATLAB 5.1) and resampled at 1024 points equally spaced on the Greenwood map. Cross-correlation functions (CCF) were calculated using the data from experiment 1 to quantify the shift between two different fine-structure patterns. The correlation lag giving the maximum of the CCF within the range ±1 mm was taken as shift between two fine-structure patterns with adjacent frequency ratios. It is assumed that small changes in frequency ratio will cause only small shifts of the overall fine structure. Therefore, the range to look for maxima of the CCF was limited to avoid ambiguities which could be caused by the quasiperiodic shape of the patterns. The computation of CCF was always restricted to the area of actual overlap between the compared patterns.
II. EXPERIMENTAL RESULTS

A. Experiment 1: Dependence on $f_2/f_1$

Figure 2 shows the effect of variation of $f_2/f_1$ (seven different $f_2/f_1$ ratios) on the DP fine-structure patterns for the $2f_1-f_2$ distortion product for four different subjects. The left column shows the results plotted as a function of $f_2$ place, and the right column shows the same data as a function of $2f_1-f_2$. The labeling of the ordinate holds for the bottom trace only. Each successive trace that corresponds to a different $f_2/f_1$ ratio is shifted by 20 dB. Note the pronounced shift in the basal direction of successive patterns when plotted as a function of $f_2$. This contrasts with the small shift in the apical direction when plotted as a function of $2f_1-f_2$. This is illustrated by the lines in the middle column showing the cumulative sum of correlation lag for the maxima in the CCF between successive patterns. The similarity between adjacent patterns is relatively high for small differences in $f_2/f_1$. However, the patterns become more different when the changes in frequency ratio get bigger.

B. Experiment 2: Fine structure of different order DPOAE

Figure 3 shows fine-structure patterns for different order DPOAE characterized by the same distance along the basilar membrane between $f_2$ and $f_{DP}$ (right column) or $f_1$ and $f_{DP}$ (left column) for the same four subjects as in Fig. 2. As Fig. 3 illustrates, the patterns for same $f_2$ and $f_{DP}$ frequencies are very similar, suggesting that the relative phase between the DP components contributing from the characteristic places of $f_2$ and $f_{DP}$ plays an important role in the DPOAE fine structure. For identical $f_1$ and $f_{DP}$ the patterns still look similar. However, for most of the subjects the CCF indicates a slight shift in the basal direction for the higher-order DPOAE. This is consistent with the movement of the $f_2$ place in the apical direction with increasing order in this case.

C. Experiment 3: Fixed $f_2$ vs fixed $f_{DP}$

While the results from experiments 1 and 2 indicate that the DPOAE observed in the human ear canal stems from two sources along the cochlea partition, one close to the $f_2$ site and one at the $f_{DP}$ site, a separation of the contribution of these two sources cannot be achieved using these data. Figure 4 shows DPOAE patterns for four subjects (two of them as in Figs. 2 and 3) obtained with a fixed $f_2$ (gray line) and with a fixed $f_{DP}$ (black line). The use of a fixed $f_2$ results in patterns very similar to the ones observed before, using the fixed-ratio paradigm. In contrast, the use of a fixed $f_{DP}$ greatly reduces the fine structure. There remains only an overall dependence of DP level on frequency ratio, with a maximum for $2f_1-f_2$ at $f_2-f_1$ around 1.2, as reported, e.g., by Harris et al. (1992).

III. SIMULATIONS IN A NONLINEAR AND ACTIVE COCHLEA MODEL

A. Description of the model

For computer simulations of the observed effects, a one-dimensional nonlinear and active model of the cochlea was used, based on a model described in previous work (van Hengel et al., 1996; van Hengel and Duifhuis, 1999). In these papers it was shown that the model, which operates in the time domain, is very useful to study nonlinear phenomena such as OAE. The basis for such models has been described in more detail in Duifhuis et al. (1985) and van den Raadt and Duifhuis (1990). In previous simulations of DPOAE, it turned out that a possible shortcoming of the model was that it did not produce a high and broad excitation peak for pure-tone stimuli (van Hengel and Duifhuis, 1999). It is claimed by various authors that such a peak is necessary to properly simulate cochlear behavior at low stimulus levels (e.g., Zweig, 1991; de Boer, 1995). Furthermore, it was claimed by Shera and Zweig (1993) and shown by Talmadge et al. (1993, 1998) that the impedance function suggested by Zweig (1991), which produces a high and broad excitation peak, also produces a fine structure in various types of simulated emissions when it is combined with a ‘‘roughness’’ in the mechanical parameters of the cochlear partition. This roughness is a random fluctuation of (one of) the parameters describing the mechanics of the sections of the cochlear partition used in the model, and reflects random inhomogeneity in the placement and behavior of cells along the cochlea, especially the outer hair cells. The impedance function described by Zweig (1991) was therefore incorporated in the model, as well as the possibility of introducing roughness. The resulting model consists of 600 sections equally spaced along the length of the cochlea (35 mm). The motion of the cochlear partition in each section is described by the following equation of motion:

\[ m \ddot{y}(x) + d(x,V) \dot{y}(x) + s(x)[y(x) + c(V)y(x)]_{t-\tau} = p(x). \]  

This is a normal second-order differential equation of motion for a harmonic oscillator with mass $m$, damping $d(x,V)$, and stiffness $s(x)$, driven by a pressure force $p(x)$ ($x$ is the position of the oscillator measured from the stapes along the cochlea, $y$ is the displacement, and $V = \dot{y}$ the velocity of the cochlear partition in the vertical direction), except that there is an additional ‘‘delayed feedback stiffness’’ term $s(x)c(V)y(x)_{t-\tau}$. This term was derived by Zweig (1991) from fits to experimental data on BM excitation patterns. It serves to stabilize the motion of the oscillator, counteracting a negative damping term $d(x,V)$. In order to do so, and to arrive at the desired high and broad peak in the excitation caused by a pure tone, the time delay $\tau$ must depend on the resonance frequency $\omega_{res} = \sqrt{s(x)/m}$ of the oscillator as $\tau = 1.742 \times 2 \pi/\omega_{res}$ (Zweig, 1991). The values of the parameters $d(x,V)$ and $c(V)$ determined by Zweig were $-0.1217 \sqrt{ms(x)}$ and 0.1416, respectively. It is important to note that these values are derived from estimated excitation patterns in the cochlea of the squirrel monkey at low levels of stimulation and in the frequency range around 8 kHz.
FIG. 2. Experiment 1: Dependence of $2f_1 - f_2$ DPOAE fine structure on the frequency ratio $f_2/f_1$ for four different subjects (from top to bottom: KI right, MG left, MK right, and MM right). Each row shows DPOAE fine-structure patterns for seven different ratios $f_2/f_1$ (from bottom to top: 1.07, 1.1, 1.13, 1.16, 1.19, 1.22). Left column: DP level as a function of $f_2$ place. 10-mm distance to base corresponds to a frequency $f_2 = 4358$ Hz, and 20 mm distance corresponds to $f_2 = 986$ Hz. Right column: same data as a function of $2f_1 - f_2$. 12 mm corresponds to $f_{DP} = 3270$ Hz, and 22 mm corresponds to $f_{DP} = 713$ Hz. The labeling of the ordinate holds for the bottom trace only. Each successive trace is shifted up by 20 dB. Middle column: shift between the different DPOAE patterns when plotted as a function of $f_2$ (black line) and when plotted as a function of $2f_1 - f_2$ (gray line). This overall shift is quantified by the correlation lag for the maximum of the cross-correlation function between adjacent patterns. Each data point results from cumulative summation of the shifts between adjacent patterns (as indicated by the numbers “1.07 to 1.1,” “1.1 to 1.13,” etc.), starting at the frequency ratio 1.07.
FIG. 3. Experiment 2: Comparison of different order DPOAE fine-structure patterns for four subjects. Left column: Identical $f_1$ and $f_{DP}$ frequencies, right column: Identical $f_2$ and $f_{DP}$ frequencies. The frequency ratios $f_2/f_1$ were chosen according to the scheme in Fig. 1. Note the very similar patterns for all orders of DPOAE. The labeling of the ordinate holds for the bottom trace only. Each successive trace is shifted by 20 dB. 13 mm corresponds to $f_{DP}=2830$ Hz. Same subjects as in Fig. 2.
These values certainly cannot be used in the vicinity of both stapes and helicotrema, since this would lead to instability (van Hengel, 1993). It is also clear that these values do not hold for higher stimulus levels. Both the negative damping term and the stabilizing delayed feedback stiffness term are thought to result from active, i.e., energy producing, behavior of the outer hair cells. This active behavior must saturate at higher levels. It is therefore logical to capture the nonlinearity of the outer hair cells. This active behavior must saturate thought to result from active, i.e., energy producing, behavior and the stabilizing delayed feedback stiffness term are hold for higher stimulus levels. Both the negative damping

\[ d(x, v) = \left[ d_1 + \frac{\beta(d_h - d_i)v}{1 + \beta|v|} \right] \frac{\sqrt{ms(x)}}{1 + \beta|v|} \]

with \( d_i = -0.12 \), \( d_h = 0.5 \),

\[ c(v) = c_l + \frac{\beta d_i v}{1 + \beta|v|} \quad c_l = 0.1416. \]

In these equations the nonlinear behavior of \( d(x, v) \) and \( c(v) \) is chosen to be the same, with the damping going to a value of \( d_h \sqrt{ms(x)} \) and the delayed feedback stiffness disappearing at high excitation levels. The region in which the nonlinearity plays a role is determined by the parameter \( \beta \). In all simulations presented here, a value of 0.01 ms/nm was used for this parameter, leading to a compressive growth of the excitation at the characteristic place for a pure-tone stimulus of around 0.3 dB/db over the range from about 20 to 80 dB SPL stimulus level. Because the mass \( m \) was chosen to be 0.375 kg/m\(^2\), independent of the position \( x \), the stiffness controls the place–frequency map of the cochlear partition. The place–frequency map chosen was

\[ f(x) = A \times e^{-ax}, \quad A = 22.508 \text{ kHz}, \quad a = 150 \text{ m}^{-1}. \]

Following the work of Shera and Zweig (1993) and Talmadge and Tubis (1993), the random fluctuations necessary to obtain a fine structure in the emissions were introduced in the stiffness as

\[ s(x) = s(x) \left[ 1 + r_0 r(x) \right], \]

where \( r(x) \) is a random variable with a Gaussian distribution and \( r_0 \) is a scaling parameter that controls the amount of roughness. For the results presented here, a value of 1% was used for \( r_0 \). The equations of motion [Eq. (1)] for all sections were coupled through the fluid, which was assumed linear, incompressible, and inviscid. The coupled system was solved by Gauss elimination and integrated in time using a Runge–Kutta 4 time-integration scheme with a sampling frequency of 150 kHz. Reducing the sampling frequency could lead to instabilities in certain cases, but increasing it did not give significantly different results (differences in emission levels were below 0.5%).

To simulate OAE, the motion of the cochlea sections is coupled to the outside world via a simplified middle ear, consisting of a mass, stiffness, and damping in combination with a transformer. This produces a sound pressure that would result at the eardrum in an open ear canal. Previous studies with this model have shown that emission levels are highly sensitive to conditions at the eardrum. SOAE level may vary up to 30 dB when different loading impedances are added (van den Raadt and Duijnhuis, 1993).

B. Simulations

All the experiments described in Sec. II (except experiment 2 for the condition of same \( f_1 \) and \( f_{DP} \) frequencies) were simulated using this computer model. In contrast to the experiments, all simulations were performed with primary levels of 50 dB SPL. The model output is the sound-pressure level at the eardrum taken from a 30-ms interval beginning 20 ms after stimulus start to avoid onset effects. The data
were analyzed using the least-squares-fit method described by Long and Talmadge (1997) to get an estimate of the spectral power of the frequency components of interest. The further analysis of pattern shifts was performed using cross correlation as described for the experimental data.

The simulation results are given in Figs. 5–7. Analogous to the experimental data in Fig. 2, Fig. 5 shows simulated DPOAE fine-structure patterns for different frequency ratios of the primaries plotted as a function of \( f_2 \) (left panel) and \( f_{DP} \) (right panel). The main experimental result, i.e., the shift of fine-structure patterns when varying the frequency ratio, is replicated very well, both qualitatively and quantitatively. Figure 6 shows fine-structure patterns for three different frequency ratios, which were chosen to get the same frequencies for \( f_2 \) and \( 2f_1-f_2, \) \( 3f_1-2f_2, \) or \( 4f_1-3f_2, \) respectively. As in the experiments, the patterns for different order DPOAE were almost identical in all these stimulus conditions. Figure 7 shows that, in the model, the fine structure disappears in a fixed \( f_{DP} \) paradigm, similar to the experimental results, while the model still produces a fine structure for fixed \( f_2. \) The only discrepancies between simulations and experimental results in the fine structure are the reduced dynamical range between maxima and minima in the model for small frequency ratios and the slightly smaller period of the frequency-dependent level variations.

Simulations have also been performed omitting the roughness in the model’s stiffness function in either (1) the frequency region above 2 kHz, or (2) below 2 kHz, or (3) with no roughness at all, to get a better understanding of the mechanism creating the fine structure in the model. DPOAE with a high frequency resolution were computed over a frequency range for \( f_2 \) from 2483 to 3084 Hz at a frequency ratio \( f_2/f_1=1.22. \) This ensured that the characteristic places of \( f_{DP} \) always fell in model sections with characteristic frequencies below 2 kHz, while the characteristic frequencies of

![FIG. 5. Computer simulation of experiment 1: Dependence of 2f_1-f_2 DPOAE fine structure on the frequency ratio f_2/f_1. Left column: DP level as a function of f_2 place. 10 mm distance to base corresponds to a frequency f_2=5027 Hz, and 16-mm distance corresponds to f_2=2045 Hz, according to the exponential place–frequency map used in the cochlea model. Right column: same data as a function of 2f_1-f_2 place. 11 mm corresponds to f_{DP} =4327 Hz, and 19 mm corresponds to f_{DP}=1304 Hz. As in Fig. 2, the shift between adjacent patterns is illustrated by the lines in the middle column of the figure based on cross correlation.](image)

![FIG. 6. Computer simulation of experiment 2: Comparison of different order DPOAE fine-structure patterns for identical f_2 and f_{DP} frequencies. The frequency ratios f_2/f_1 were chosen according to the scheme in Fig. 1. Note the very similar patterns for all orders of DPOAE. The labeling of the ordinate holds for the bottom trace only. Each successive trace is shifted by 20 dB. 14 mm corresponds to f_{DP}=2760 Hz, according to the exponential frequency map used in the model.](image)

![FIG. 7. Computer simulation of experiment 3: DP level as a function of f_1. Black line: fixed f_{DP} paradigm, i.e., varying both f_1 and f_2 while keeping f_{DP} fixed at 2 kHz. Gray line: fixed f_2 paradigm, i.e., varying f_1 and f_{DP} while f_2 is fixed at 3 kHz. Note that, as for the experimental results, the fine structure for the fixed-f_2 paradigm is similar to the patterns observable with the fixed-ratio paradigm, while there is no fine structure when using the fixed-f_{DP} paradigm.](image)
the primaries always fell into sections above 2 kHz. Figure 8 shows the simulation results in these three conditions as well as in the reference condition with roughness over the whole length of the cochlea. The DPOAE fine structure is unaffected by the presence or absence of the roughness in the primary region, while it disappears when the roughness is omitted in the distortion product frequency region. This emphasizes the interpretation of the experimental results that the DPOAE fine structure is mainly influenced by the re-emission components from the characteristic DP places, while emission from the primary component places is almost constant in level and phase over frequency.

IV. DISCUSSION

Similar to recent experimental and theoretical studies on DPOAE fine structure by other authors (e.g., Mauermann et al., 1997a; Heitmann et al., 1998; Talmadge et al., 1998, 1999), our experiments and simulations give further evidence that the fine structure is the result of two sources. Furthermore, the idea is supported that the underlying physical mechanisms of these two sources are different or at least act in a different way (e.g., Shera and Guinan, 1999).

To illustrate this, the results presented in this paper will be interpreted in three steps. The results of experiment 1 show that the DPOAE fine structure is not caused by local mechanical properties of the primary region, but rather that the characteristic site of the distortion product frequency plays a crucial role. Experiment 2 shows that independent of the distortion product order, fine-structure patterns are very similar as long as the characteristic sites of \( f_2 \) and of the DP frequencies are the same, i.e., the relative phase between the two emission sites is almost constant in this experiment. Taken together with the findings from experiment 1, this implies that the emission recorded in the ear canal is the vector sum of components from these two sites. The relative phase of these components has at least some influence on the DPOAE fine-structure pattern.

The results of experiment 3 show a quasiperiodic variation when the re-emission site is varied in frequency monotonically while the generation site is held fixed, whereas the quasiperiodic variation in DP level disappears in the case of a fixed re-emission site and a sweep of the \( f_2 \) frequency. In terms of a vector summation of two components, this indicates that the component generated at the primary place must be almost constant in level and phase (at least locally) regardless of the frequency. To explain the quasiperiodic variations in the sum of the two components, we have to assume that the re-emission component varies either in phase or in level with increasing frequency. Shera and Zweig (1993) showed that the fine structure of stimulus frequency otoacoustic emissions (SFOAE) can be interpreted as interference of the incoming and outgoing traveling waves with a periodic rotating phase of the cochlear reflectance. It appears reasonable to treat the DPOAE re-emission component in a similar way to SFOAE generation. Therefore, a periodically varying phase of the re-emission component is the most likely explanation for the DPOAE fine structure. Following, furthermore, the arguments of the “Gedankenexperiment” described by Shera and Guinan (1998), the different characteristics of the two components (rotating phase of the \( f_\text{DP} \) component, almost constant phase of the \( f_2 \) component) indicate that the underlying mechanisms for these two DPOAE components are different. These authors (Shera and Guinan, 1998, 1999) distinguished two classes of OAE mechanisms, “linear coherent reflection” and “nonlinear distortion,” whereby DPOAE are a combination of both a nonlinear distortion at the generation site and a coherent reflection from the characteristic site of \( f_\text{DP} \).

The disappearance of fine structure during the fixed \( f_\text{DP} \) experiment shows that there is no coherent reflection from the primary region because there is no rotating phase with frequency. Using the approach suggested by Zweig and Shera (1995) to explain the spectral periodicity of reflection emissions, this had to be expected because the traveling wave from the initial generation site results in constant wavelength only around the \( f_\text{DP} \) site but not in the region around the primaries. Therefore, the contribution from the primary region needs to be generated in a different way.
most probably due to nonlinear distortion. This interpretation of the experiments is confirmed by the computer simulations, which show no effect on the fine structure when removing the roughness (which is necessary for coherent reflections) from the primary region, while the fine structure disappears in simulations when removing the roughness only around the \( f_{DP} \) region (see Fig. 8).

The overall good correspondence between simulations and experimental results gives further confirmation for a whole class of two-source interference models like the one used here. This class of models was recently described in detail by Talmadge et al. (1998). The simulations with partly removed roughness (see Fig. 8) cannot directly be transformed into an experimental approach because it is impossible to “flatten out” certain areas of the human cochlea. But, a situation close to that might be studied. If a local area of the cochlea is damaged, most probably no broad and tall excitation pattern can be built up there, which in addition to roughness is necessary for coherent reflections (Zweig and Shera, 1995; Talmadge et al., 1999). To get further insight into the mechanisms of DPOAE fine structure, this approach is investigated in the accompanying paper (Mauermann et al., 1999) by looking at the DPOAE fine structure of subjects with frequency-specific hearing losses.

V. CONCLUSIONS

Distortion products recorded in the ear canal cannot be traced back to one single source on the basilar membrane. Instead, DPOAE fine structure reflects the interaction of two components with different underlying physical principles. The first component is due to nonlinear distortion at the primary site close to \( f_2 \) and has a nearly constant phase and level. The second component is caused by a coherent reflection from the re-emission site at the characteristic place of \( f_{DP} \) and shows a periodically varying amplitude or phase when changing the primary frequencies.

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1The number of sections had to be increased from the original 400 to at least 600 in order to avoid “wiggles” in the excitation patterns. These “wiggles” were also found by Talmadge and Tubis in their work on a time-domain model involving the “Zweig-impedance” and made them use a spatial discretisation of 4000 sections (Talmadge and Tubis, 1993; and personal communication).

2Of course, other terms could also contain nonlinearity. For example, Furst and Goldstein (1982) argue that the stiffness term should be made nonlinear. For reasons of simplicity only \( d(x,v) \) and \( c(v) \) were made nonlinear here, since these two terms must certainly change with input level.

3There is another discrepancy between the experimental results and the simulations in the overall shape of the patterns. For the range of frequency ratios observed here, we see a maximum in DPOAE level for human subjects at a frequency ratio around 1.225 (Gaskill and Brown, 1990) while the level is reduced for smaller and larger frequency ratios. This reflects the so-called “second filter” effect (e.g. Brown and Williams, 1993; Allen and Fahey, 1993), which is currently not included in our simulations. With the parameter settings used in this study, the second filter effect produced by the model does not resemble the shapes found in the experimental data. However, as described in van Hengel and Duifhuis (1999), the second filter behavior can also be simulated using this kind of transmission-line model. Therefore, in the near future attempts will be made to improve the model results by finding parameter values fitting both fine structure and the second filter.


