Neural encoding of voice pitch and formant structure at birth as revealed by frequency-following responses

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Detailed neural encoding of voice pitch and formant structure plays a crucial role in speech perception, and is of key importance for an appropriate acquisition of the phonetic repertoire in infants since birth. However, the extent to what newborns are capable of extracting pitch and formant structure information from the temporal envelope and the temporal fine structure of speech sounds, respectively, remains unclear. Here, we recorded the frequency-following response (FFR) elicited by a novel two-vowel, rising-pitch-ending stimulus to simultaneously characterize voice pitch and formant structure encoding accuracy in a sample of neonates and adults. Data revealed that newborns tracked changes in voice pitch reliably and no differently than adults, but exhibited weaker signatures of formant structure encoding, particularly at higher formant frequency ranges. Thus, our results indicate a well-developed encoding of voice pitch at birth, while formant structure representation is maturing in a frequency-dependent manner. Furthermore, we demonstrate the feasibility to assess voice pitch and formant structure encoding within clinical evaluation times in a hospital setting, and suggest the possibility to use this novel stimulus as a tool for longitudinal developmental studies of the auditory system.

Spoken language is arguably the most prevalent form of human communication. Experimental evidence suggests a universal organic basis for language acquisition, based on the identical development of speech perception pathways observed across different populations, languages and cultures1–3. Speech perceptual skills have been well characterized along the lifespan, especially with regard to their maturation during the first year4–6. However, less is known about their functional state during the very first hours after birth, when humans newly encounter the rich and challenging complexity of the external acoustic environment. A highly efficient auditory system becomes hence a requisite for proper language acquisition, as the complex and dynamic acoustic signal of speech conveys only very slight spectral and temporal cues for speech sound discrimination7.

Previous studies have shown that the auditory system of newborns and infants is able to handle several aspects related to pitch processing, such as tracking pitch contours8–15, higher-order frequency direction relationships16, processing a missing fundamental17 or exhibiting relative pitch by discriminating transposed melodies18,19. Likewise, the newborn auditory system appears able to discriminate phonemes20–22 even when only based upon vowel formant structure changes or duration20,23,24. And yet, the low level neural underpinning of these abilities remains to be established.

Using non-invasive electroencephalography recordings, auditory brainstem responses (ABR) evoked to acoustic transients, such as click stimuli, have successfully been used to assess the integrity of the auditory pathway25–27. However, periodic acoustic stimuli also elicit a particular brain response of subcortical and cortical origin, known as the frequency-following response (FFR)28–30. The FFR reflects with high fidelity the encoding of periodic temporal envelope modulations (FFRENV) and temporal fine structure harmonic constituents (FFRTFS) of a stimulus (Aiken and Picton31 following the terminology proposed by Krizman and Kraus29). In language studies, these

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two components of the FFR have been respectively regarded as indexes of two perceptual properties of speech sounds: voice pitch contour and formant structure.

FFR recordings are increasingly considered a valuable tool to index the current functional state of the auditory system and to predict the future development of language, since disruptions in the FFR elicited by speech sounds relate to deficits in phonological awareness, reading impairments and dyslexia. The potential of the FFR as a biomarker for auditory deficits and their relation to literacy skills has thus been proposed. However, most developmental studies on the FFR targeted babies of several months of age, toddlers, infants or years-old children, with only a few published reports on newborns. Thus, knowledge about the expected speech perceptual skills in newborns, who are more vulnerable than older age groups to hearing damage, may aid the early detection of language impairments and guide appropriate interventions benefitting from the massive neural plasticity during the first years of life.

Moreover, while newborn studies have focused on the assessment and maturation of pitch processing through the analysis of the FFR env, to date, none addressed formant structure encoding in a systematic manner. To the best of our knowledge, only a recent study from our lab providing a normative newborn database of FFR env, properties, attempted, as a secondary aim, to reveal whether the neonate auditory system was able to discriminate sounds differing in their fine structure (/da/ vs. /ga/). While the results were negative, the apparent lack of formant structure encoding may be due to several reasons: (a) the short duration of the consonant transition (47 ms) and its formant change, which limit the resolution of the computed spectral information; (b) the high frequency content of the stimuli (/da/ vs. /ga/: F0 = 1438–1214 Hz, /ga/ = 1801–1214 Hz), which elicits diminished FFR amplitudes that are difficult to recognize; (c) the fact that phase-locking to higher frequencies develops later than to lower ones; and, ultimately, (d) the nature of the analyzed signal (FFR env), which emphasizes temporal envelope information representation at the expense of temporal fine structure. Thus, it still remains unclear whether newborns cannot yet precisely track formant changes in complex sounds or if stimulation parameters used so far were not suited to reveal this ability.

Furthermore, in newborn FFR research, time is of the essence. Recording time constraints determine what stimulus encoding abilities can be studied, and it is even a more challenging issue when newborn research is conducted in a hospital setting. In addition to the ease of waking up or disturbing sleep, the hospital environment requires frequent and continuous access to the baby and the mother for routine tests to discard serious health issues, interventions, in-depth health evaluations, and any other kind of neonatal care. Taking this into account, it would be unsuitable to carry out recording sessions lasting between 40 and 50 min, typical of speech-sound discrimination studies. The most adequate session duration would be between 20 and 30 min. Therefore, devising a new single stimulus that would allow a proper assessment of both the FFR env and FFR tsp simultaneously within recording times compatible with infant research is required. This would provide a snapshot of the functional state of speech sound processing mechanisms at birth and, ultimately, help better understand how the encoding of this complex auditory signal matures.

Thus, the aim of the present study was to characterize the functional maturity of voice pitch contour and formant structure encoding mechanisms in the newborn population with non-invasive electrophysiological recordings, using the adult population as a reference. To that end, we developed a novel speech stimulus that allows the simultaneous assessment of both components of the speech signal through analyzing the FFR env and the FFR tsp, and which is at the same time compatible with clinical evaluation time constrains in a hospital setting. FFRs were recorded to a novel two-vowel (/oa/’/ vowel) speech stimulus with a rising pitch ending. In order to estimate voice pitch encoding from the temporal envelope of the recorded neural response, we computed the FFR env and analyzed it (spectral measures at the fundamental frequency [F0] peak; pitch tracking measures extracted from stimulus-to-response cross-correlations and signal autocorrelations) accounting for the different steady and rising pitch sections of the stimulus (both during the /a/ vowel). In order to estimate formant structure encoding from the temporal fine structure of the recorded response, we computed the FFR tsp and analyzed it (spectral measures at the first formant [F1] frequency peak) accounting for the different /o/ and /a/ vowel sections of the stimulus (both during the steady pitch section).

Given that the human auditory system is able to encode changes in voice pitch with great precision starting from the first days of life, we would expect no significant differences between newborns and adults in spectral amplitudes at F0 or in pitch tracking accuracy measures computed from the FFR env. Because the capacity of neurons in the auditory system to phase-lock their activity to higher sound frequencies develops later than to lower ones and continues improving during the first year of life, we would expect newborns to exhibit overall smaller FFR tsp spectral amplitudes at F1 peak frequencies than adults. However, we had no clear hypotheses as to whether the newborn FFR tsp would reflect a discriminative encoding of vowel formant structure as it does in adults and, if so, whether that discriminative encoding would depend on F1 center frequency. The evidence suggesting that the newborn auditory system discriminates vowel changes, is based upon recordings of event-related potentials (ERPs) reflecting higher-order auditory system computations. But, the fundamentally different nature of the FFR as a phase-locked neural response reflecting the acoustic waveform with high precision, and the fact that this low-level encoding of acoustic features is immature at birth, at least for higher frequencies, precluded us at this point to have strong hypotheses on the newborn’s auditory system’s capabilities.

**Results**

Temporal envelope-following responses (FFR env) and temporal fine structure-following responses (FFR tsp) elicited by a two-vowel syllable, rising-pitch ending, /oa/ stimulus (Fig. 1a) were collected from 34 newborns and 18 adult participants. In order to assess voice pitch contour and formant structure encoding in depth, neural responses were analyzed according to the sound features of the different stimulus sections. Below, we provide descriptive statistics and comparisons for a comprehensive number of parameters extracted from the FFR env.
and FFRTFS (see “Methods” for a detailed description). Statistically non-significant results can be found in Suppl. Table 1.

**Figure 1.** Temporal representation of the stimulus (a); FFR\textsubscript{ENV} (b) and FFR\textsubscript{TFS} (c). (a) Time waveform (top) and spectrogram of the /oa/ stimulus with schematic overlay of the formant structure trajectory (targeted F\textsubscript{0} and F\textsubscript{1} in solid lines; non-analyzed F\textsubscript{2} depicted in dotted line). (b) Grand averaged time-domain waveform of the FFR\textsubscript{ENV} from newborns (top red) and adults (bottom blue), obtained by averaging the neural responses to the two stimulus polarities. (c) Grand averaged time-domain waveform of the FFR\textsubscript{TFS} from newborns (top red) and adults (bottom blue), obtained by subtracting the neural responses to the two stimulus polarities.
Neural transmission delay. Neural lag. Newborns showed a significantly longer neural lag (an estimation of FFR latency) compared to adults ($U_{(50)} = 59$, $p < 0.001$, Cohen’s $d = 0.659$). Descriptive statistics can be found in Table 1.

Assessment of voice pitch encoding from FFR\textsubscript{ENV}. In order to determine the strength of the representation of the $F_0$ and assess the accuracy in tracking $F_0$ changes, our /oa/ stimulus was devised to feature a steady pitch during its initial section (113 Hz; 0–160 ms) and a linearly increasing pitch during its final section (113–154 Hz; 160–250 ms) (Fig. 1a). To accentuate the FFR components corresponding to the encoding of the stimulus envelope (mainly the $F_0$) while suppressing those related to the fine structure, thus controlling for vowel changes that occur along the different sections of the stimulus, we computed the FFR\textsubscript{ENV}. Grand-average FFR\textsubscript{ENV} waveforms are shown in Fig. 1b for both groups separately (newborns and adults). All descriptive statistics for FFR\textsubscript{ENV} derived parameters can be found in Table 1.

Spectral amplitude at $F_0$ peak. The spectral amplitude at $F_0$ peak (113 Hz) during the steady pitch section of the stimulus (10–160 ms) was calculated as an indicator of the magnitude of neural phase-locking at that specific frequency. Newborns exhibited significantly reduced spectral amplitudes at $F_0$ peak as compared to adults ($t_{(50)} = − 3.079$, $p = 0.003$, Cohen’s $d = − 0.831$). The corresponding amplitude spectra in the frequency domain computed along the steady pitch stimulus section is shown in Fig. 2a. Figure 2b illustrates the distribution of $F_0$ spectral amplitude values obtained for each group.

Signal-to-noise ratio. The signal-to-noise ratio (SNR) at $F_0$ peak during the steady pitch section of the stimulus was taken as an estimation of the relative spectral magnitude of the response. No significant group differences were found. Figure 2c illustrates the distribution of $F_0$ SNR values obtained per group.

### Table 1. Descriptive statistics for FFR\textsubscript{ENV} derived parameters: neural lag; $F_0$ spectral amplitude and SNR computed for the steady pitch section; stimulus-to-response cross-correlation, pitch error and pitch strength computed separately for each section of the stimulus (/a/ steady section; /a/ rising section). SD standard deviation, $Q_1$ first quartile (25th percentile), $Q_3$ third quartile (75th percentile), IQR interquartile range.

| Measure                        | Mean  | SD   | Median | $Q_1$ | $Q_3$ | IQR  | Minimum | Maximum |
|-------------------------------|-------|------|--------|-------|-------|-------|---------|---------|
| Neural lag (ms; from 10 to 250 ms) |       |      |        |       |       |       |         |         |
| Newborns                      | 9.33  | 1.74 | 9.26   | 8.70  | 10.01 | 1.31  | 3.53    | 12.60   |
| Adults                        | 6.26  | 1.21 | 5.85   | 5.55  | 6.47  | 0.92  | 5.10    | 9.75    |
| $F_0$ spectral amplitude ($\mu$V; from 10 to 160 ms) |       |      |        |       |       |       |         |         |
| Newborns                      | 0.01  | 0.01 | 0.01   | <0.01 | 0.02  | 0.01  | <0.01   | 0.03    |
| Adults                        | 0.02  | 0.01 | 0.02   | 0.01  | 0.02  | 0.01  | 0.01    | 0.04    |
| $F_0$ SNR (from 10 to 160 ms) |       |      |        |       |       |       |         |         |
| Newborns                      | 4.36  | 4.72 | 5.86   | 0.82  | 7.83  | 7.01  | −11.07  | 10.95   |
| Adults                        | 4.53  | 3.43 | 4.70   | 3.01  | 7.26  | 4.25  | −3.52   | 9.40    |
| /a/ steady section (90–160 ms) |       |      |        |       |       |       |         |         |
| Cross-correlation (Pearson’s r)|       |      |        |       |       |       |         |         |
| Newborns                      | 0.18  | 0.06 | 0.18   | 0.12  | 0.23  | 0.11  | 0.06    | 0.30    |
| Adults                        | 0.20  | 0.05 | 0.20   | 0.17  | 0.23  | 0.07  | 0.07    | 0.27    |
| Pitch error (Hz)              |       |      |        |       |       |       |         |         |
| Newborns                      | 11.66 | 7.15 | 10.18  | 5.64  | 16.55 | 10.90 | 2.76    | 28.73   |
| Adults                        | 9.45  | 3.40 | 9.77   | 6.45  | 12.07 | 5.63  | 3.27    | 15.29   |
|Pitch strength (r)             |       |      |        |       |       |       |         |         |
| Newborns                      | 0.60  | 0.18 | 0.57   | 0.45  | 0.74  | 0.29  | 0.32    | 0.88    |
| Adults                        | 0.55  | 0.09 | 0.55   | 0.46  | 0.62  | 0.15  | 0.43    | 0.76    |
| /a/ rising section (160–250 ms) |       |      |        |       |       |       |         |         |
| Cross-correlation (Pearson’s r)|       |      |        |       |       |       |         |         |
| Newborns                      | 0.11  | 0.04 | 0.10   | 0.08  | 0.13  | 0.05  | 0.05    | 0.18    |
| Adults                        | 0.10  | 0.02 | 0.10   | 0.09  | 0.12  | 0.03  | 0.06    | 0.15    |
| Pitch error (Hz)              |       |      |        |       |       |       |         |         |
| Newborns                      | 11.72 | 7.08 | 10.44  | 5.50  | 16.25 | 10.75 | 2.60    | 28.15   |
| Adults                        | 9.50  | 3.34 | 9.87   | 6.55  | 12.00 | 5.45  | 3.13    | 15.03   |
|Pitch strength (r)             |       |      |        |       |       |       |         |         |
| Newborns                      | 0.60  | 0.18 | 0.56   | 0.45  | 0.74  | 0.29  | 0.32    | 0.88    |
| Adults                        | 0.55  | 0.09 | 0.55   | 0.46  | 0.61  | 0.15  | 0.44    | 0.75    |
Stimulus-to-response cross-correlation. The stimulus-to-response cross-correlation was taken as a measure of the accuracy with which the FFR_{ENV} reproduced the stimulus waveform, separately for the /a/ steady and /a/ rising pitch contour stimulus sections. Lower stimulus-to-response cross-correlation values were obtained during the rising pitch section (mean ± SD; /a/ rising = 0.11 ± 0.03) as compared to the steady pitch section (mean ± SD; /a/ steady = 0.18 ± 0.06) (Z = −5.774, \(p < 0.001\), Cohen's \(d = 0.801\)). No significant group differences or group per stimulus section interaction were found.

Pitch error. We then computed the pitch error per pitch section separately, in order to determine pitch-tracking accuracy of the F0 contour\(^{11,29}\). Neither significant group or stimulus section differences nor group per stimulus section interaction were found (see Fig. 3a for spectrogram and Fig. 3b for pitch track).

Pitch strength. Pitch strength was taken as a measure of periodicity and the magnitude of neural phase-locking of the response\(^{10}\) and was also computed separately per stimulus pitch section. Neither significant group or stimulus section differences nor group per stimulus section interaction were found.

Assessment of formant structure encoding from FFR_{TPS}. In order to determine the ability of the participants to encode the formant structure of speech sounds, the /oa/ stimulus featured two sections with steady pitch but differing in their formant structure: the /o/ section (10–80 ms; \(F_1 = 452\) Hz) and the /a/ steady pitch section (90–160 ms; \(F_1 = 678\) Hz). In order to emphasize temporal fine structure components of the response while diminishing the contribution of responses to the temporal envelope, we computed the FFR_{TPS}\(^{29,31}\). Grand-average FFR_{TPS} waveforms are shown in Fig. 1c for both groups separately. The frequency spectrum of the /o/ section and the /a/ steady pitch section are shown in Fig. 4a for both groups. All descriptive statistics can be found in Table 2.

Spectral amplitudes and SNRs from the FFR_{TPS} were retrieved separately from neural responses during the /o/ section (10–80 ms) and the /a/ steady pitch section (90–160 ms), selecting the spectral peaks corresponding to stimulus \(F_1\) frequencies (452 Hz [/o/] and 678 Hz [/a/]), as indicators of the magnitude (absolute and relative) of phase-locking at the selected frequencies.

Spectral amplitude at /o/ vowel \(F_1\). Spectral amplitudes at the /o/ vowel \(F_1\) (452 Hz) are illustrated in Fig. 4b (left). A main effect of group revealed significantly smaller spectral amplitudes at 452 Hz in newborns as compared to adults (\(F_{(1,50)} = 85.778\), \(p < 0.001\), \(\eta^2 = 0.632\)). A main effect of stimulus section showed a significantly larger spectral amplitude value at 452 Hz during the /o/ vs. /a/ steady pitch sections (\(F_{(1,50)} = 25.529\), \(p < 0.001\), \(\eta^2 = 0.338\)). The group per stimulus section interaction was significant as well (\(F_{(1,50)} = 18.603\), \(p < 0.001\), \(\eta^2 = 0.271\)). Post-hoc tests computed to determine the direction of the interaction revealed higher spectral...
Spectral amplitudes in adults at 452 Hz during the /o/ vs. /a/ sections ($t(17) = 3.803$, $p = 0.001$, Cohen's $d = 0.896$), but no significant differences were found in newborns.

Spectral amplitude at /a/ vowel $F_1$. Spectral amplitudes at the /a/ vowel $F_1$ (678 Hz) are illustrated in Fig. 4b (right). A main effect of group revealed significantly smaller spectral amplitudes at 678 Hz in newborns as compared to adults ($F(1,50) = 79.157$, $p < 0.001$, $\eta^2 = 0.613$). A main effect of stimulus section showed a significantly larger spectral amplitude value at 678 Hz during the /a/ steady pitch vs. /o/ sections ($F(1,50) = 64.555$, $p < 0.001$, $\eta^2 = 0.564$). The group per stimulus section interaction was significant as well ($F(1,50) = 50.252$, $p < 0.001$, $\eta^2 = 0.501$). Post-hoc tests computed to determine the direction of the interaction revealed higher spectral amplitudes in adults at 678 Hz during the /a/ steady pitch vs. /o/ sections ($t(17) = -5.845$, $p < 0.001$, Cohen's $d = -1.378$), but no significant differences were found in newborns.

SNR at /o/ vowel $F_1$. SNR values at the /o/ vowel $F_1$ (452 Hz) are illustrated in Fig. 4c (left). A main effect of group revealed significantly smaller SNR values at 452 Hz in newborns as compared to adults ($F(1,50) = 47.213$, $p < 0.001$, $\eta^2 = 0.486$). A main effect of stimulus section showed a significantly larger SNR value at 452 Hz during the /o/ vs. /a/ steady pitch sections ($F(1,50) = 4.207$, $p = 0.046$, $\eta^2 = 0.078$). No significant group per stimulus section interaction was found.

SNR at /a/ vowel $F_1$. SNR values at the /a/ vowel $F_1$ (678 Hz) are illustrated in Fig. 4c (right). A main effect of group revealed significantly smaller SNR values at 678 Hz in newborns as compared to adults ($F(1,50) = 17.136$, $p < 0.001$, $\eta^2 = 0.235$). A main effect of stimulus section showed a significantly larger SNR value at 678 Hz during the /a/ steady pitch vs. /o/ sections ($F(1,50) = 15.414$, $p < 0.001$, $\eta^2 = 0.236$). The group per stimulus section interaction was significant as well ($F(1,50) = 4.753$, $p = 0.034$, $\eta^2 = 0.087$). Post-hoc tests computed to determine
the direction of the interaction revealed higher SNR values in adults at 678 Hz during the /a/ steady pitch vs. /o/ sections \((t(17) = -5.656, p < 0.001, \text{Cohen's } d = -1.333)\), but no significant differences were found in newborns.

It should be noted that some SNR values, especially those of newborns at 678 Hz peak, were very close to zero. In order to ascertain whether there was a measurable signal when expected (at 452 Hz during the /o/ section and at 678 Hz during the /a/ section), we submitted the SNR values, per group and per condition separately, to one-tailed, one sample t-tests against zero. Results demonstrated that newborns had a measurable signal for lower

![Figure 4. Formant structure encoding in newborns and adults. (a) Amplitude FFR_{TFS} spectra extracted from the /o/ vowel section (green) and the /a/ vowel section (orange) from the stimulus, plotted separately for newborns (top) and adults (bottom). (b) Main effects graphic of F1 spectral amplitude at 452 Hz (left) and 678 Hz (right) during the /o/ vowel section and the /a/ vowel section, plotted in red and blue lines for newborns and adults, respectively. (c) Main effects of F1 SNR at 452 Hz (left) and 678 Hz (right) during the /o/ vowel section and the /a/ vowel section, depicting neural responses from newborns (red) and adults (blue).](image-url)
frequency formants, as shown by significant differences in SNR at 452 Hz during the /o/ section \( (t(33) = 2.407, p = 0.022, \text{Cohen's } d = 0.414) \), but no clear response at 678 Hz during the /a/ section \( (p = 0.602) \). Intriguingly, the SNR value at 678 Hz during the /o/ section was negative and significantly different from zero (mean ± SD: \( -2.08 ± 4.80 \text{ dB} \); \( t(33) = -2.530, p = 0.016, \text{Cohen's } d = -0.433 \)). Adult participants exhibited a measurable signal in the two conditions: at the 452 Hz peak during the /o/ section \( (t(17) = 17.737, p < 0.001, \text{Cohen's } d = 4.194) \) and at the 678 Hz during the /a/ section \( (t(17) = 19.043, p < 0.001, \text{Cohen's } d = 4.491) \).

### Discussion

We hereby provide an in-depth characterization of the neural encoding of speech sound features that newborns exhibit during their first hours of life, by comparing FFRs from healthy newborns and normal-hearing adult participants elicited by a novel, two-vowel /oa/ stimulus, with a rising pitch ending. Regarding the FFR parameters indexing voice pitch encoding, extracted from the FFRENV, our results support previous findings showing no significant differences in voice pitch encoding ability at birth as compared to adults, as can be appreciated from the SNR values at F0 peak as well as in pitch tracking measures, such as stimulus-to-response cross-correlation, pitch error and pitch strength. Concerning the FFR parameters indexing formant structure encoding, extracted from the FFRTFS, as expected, newborns exhibited overall diminished amplitudes than adults at both F1 peaks of interest (452 and 678 Hz). On the other hand, obtained SNR values in newborns were higher at 452 Hz (/o/ F1) during the /o/ section than during the /a/ section but not different at 678 Hz (/a/ F1), revealing the functional state of formant structure encoding mechanisms, which appear to be partially developed but still to mature, especially at higher frequency ranges. Furthermore, our results prove the feasibility to record and assess simultaneously both voice pitch and formant structure encoding within a thirty-minute period, a time-span compatible with clinical settings that allows obtaining the FFRENV and the FFRTFS in large samples of newborns.

### Considerations on the mother’s womb acting as an acoustic filter and speech perceptual skills at birth.

Speech perception abilities are crucial for early phonetic discrimination\(^{1,2,5,6}\). Human hearing begins approximately at the 26th week of fetal life and most of the development takes place between the 26th and 28th week of gestation\(^{26,39-42}\), when hair cells and their connections to the cochlea are mature enough to tune in to specific frequencies. In this regard, previous research showed that fetuses can hear and remember language sounds and may learn about several sound properties while in the womb\(^{61}\). Studies in newborns have shown a preference for their mother’s voice\(^{65}\) and for their native language\(^{46,47}\), as well as behavioral recognition of children’s stories heard only during pregnancy\(^{66}\). But, what speech sound features do babies rely upon to exhibit such identification skills? Considering that the mother’s womb acts as a low-pass filter, the sounds...
available to a fetus during the gestation period are dominated by a low frequency content (<500 Hz), while higher frequency ranges, which characterize most of the temporal fine structure of speech, would only be fully available at birth. Indeed, neonates may base their preferences on pitch contours and slow temporal dynamics, features available during pregnancy. Furthermore, albeit previous studies have shown neural signatures of vowel change detection for vowel pairs differing only in second formant (F2) frequencies in newborns and 6-months-old babies, recent electrophysiological and behavioral evidence suggests that infant vowel discrimination relies more strongly on F1 (usually below 800 Hz) than F2 frequency differences. For instance, in a comprehensive study, McCarthy et al. analyzed neural responses to vowel changes using all pairs of a set of 7 English vowels, and showed that phonetic development from 4 to 11 months-old exhibits an increasing sensitivity to higher-frequency acoustic information (i.e., infants progressively rely less on F1 changes and more on F2 changes). Importantly, while youngest infants (4–5 months-old) neural responses appeared to reflect vowel acoustics (i.e., larger acoustic changes were reflected by larger neural response changes), those from older infants (10–11 months-old) seemed to represent putatively categorical changes (i.e., vowel space maps recreated from neural data showed large differences between vowel pairs with small acoustic differences). Intriguingly, a close inspection of their data (particularly at Fig. 4) strongly suggests that vowel pairs with lower F1 frequency content (/i/ vs. /u/; < 500 Hz) are represented in youngest infants' vowel space farther apart from each other than vowel pairs with higher F1 frequency content (/a/ vs. /ɛ/; > 500 Hz), a pattern not apparent in older infants. However, the authors did not explicitly test this hypothesis. In fact, to the best of our knowledge, there is no behavioral or neurophysiological study in newborns or young infants explicitly testing vowel discrimination as a function of formant frequency. This may constitute an exciting avenue for future research linking auditory neural responses to auditory pathway and vowel discrimination development.

Regarding our data, in view of the above and taking into account that 1) the chosen first formants of our stimulus fell below (/o/ F1) and above (/a/ F1) the 500 Hz filter cut-off; 2) FFR spectral amplitudes increase with age; 3) FFR spectral amplitudes diminish along the frequency axis; and 4) FFRs are plastically modulated by experience, it appears reasonable to expect certain degree of response in newborns at the lower frequency formant (452 Hz) and a fast decay of spectral power at the higher frequency formant (678 Hz). In any case, it seems plausible that certain speech sound processing skills were already mature at birth due to a greater exposure during pregnancy, while others would still be undeveloped.

**Functional maturity state differences across speech perceptual skills at birth.** A first indicator of auditory system's functional maturity is auditory transmission delay. Measuring wave V latencies and stimulus-to-response neural lags (which were consistent with activity generated in the brainstem) we found, in agreement with previous literature, shortened delays in adult participants, which may be due to the increasing myelination and age-related changes in synaptic function. However, even with a still maturing transmission speed, our results demonstrate that newborns accurately encode the F1 of speech sounds as well as track changes in voice pitch during immediate postnatal hours, in line with previous studies. Although spectral amplitudes at the F2 peak were smaller in newborns as compared to adults, no significant differences were found with the adult sample when choosing relative amplitude measurements (i.e., SNR). Thus, the higher spectral amplitude values for adults could be due to the fact that, even during the pre-stimulus period, they also presented a higher spectral noise level (pre-stimulus root mean square: newborns = 0.03 ± 0.01 µV; adults = 0.05 ± 0.02 µV; U[51] = 571, p < 0.001).

On the other hand, our results indicate a differential processing of formant structure in newborns in comparison to adults. Similar to the results on the FFRTFS, neonates showed significantly smaller FFRTFS absolute spectral amplitude values, but also smaller relative measures such as the SNR. However, our data demonstrate that newborns can encode the fine structure of speech sounds at higher frequency ranges, as evidenced by the fact that their SNR values were higher at 452 Hz (/o/ F1) during the /o/ section than during the /a/ section, but at 678 Hz (/a/ F1) they were not significantly different from zero. Although the SNR at 678 Hz during the /o/ section was negative in newborns, when analyzing the amplitude of the frequency spectrum (Fig. 4a) we observed that spectral amplitudes at 678 Hz during either of the two sections were very weak. Because of the reduced spectral amplitude and its large standard deviation, we considered this negative value as negligible, probably due to a noisy signal at higher frequencies rather than to active inhibition.

We considered the possibility that our results regarding formant structure encoding could be influenced by the internal structure of the stimulus, i.e., the /o/ section always preceded the /a/ section. As infants and neonates seem to preferentially use rhythmic cues to segment syllables and words from the acoustic stream, newborns may be more sensitive to sound onsets than codas. According to the temporal sampling framework hypotheses, put forward by Goswami, rhythmic amplitude envelope modulations would entrain cortical oscillatory activity to exert a preferential processing of syllable onsets. However, there is no obvious reason why such preferential onset processing should be apparent only at formant structure encoding and not at pitch encoding. Therefore, in order to shed some light on this possible confounding factor, we decided to statistically compare the SNR values at F1 during the /o/ steady pitch section (10–80 ms) vs. the /a/ steady pitch section (90–160 ms), using a paired-samples t-test for each group of age. Our results showed that there were no significant differences in the SNR values at F1 between stimulus steady pitch sections for either of the two groups (newborns: t[33] = −1.466, p = 0.152; Cohen’s d = −0.251; adults: t[17] = 0.797, p = 0.436, Cohen’s d = 0.188; for further statistical information, the reader is referred to Suppl. Table 1). Thus, no onset effect in pitch encoding was observable in any group. Moreover, given the rhythmic stimulation used in our study (SOA = 295 ms), half cycle of an entrained oscillation would last enough to cover, with the high excitability phase, both /o/ and /a/ steady pitch sections of our stimulus. Furthermore, the high frequency ranges we are dealing with in our FFR data (beyond 100 Hz) are more prone to elicit recordable subcortical activity than cortical, and the modulation of phase-locking in
subcortical neuronal ensembles by cortical oscillations has not been described, to the best of our knowledge, in the literature. Finally, in our study, the adult FFR
\textsubscript{ENV} SNR values at the formant peaks showed a double dissociation, being larger at the /o/ F\textsubscript{1} frequency during the /o/ section and at the /a/ F\textsubscript{1} frequency during the /a/ section, ruling out any onset effect. Therefore, given the pattern of results and the reviewed literature, an onset effect seems a negligible influencing factor in our results. In any case, further research studying the influence of vowel order should be carried out to help better clarify this possible confound (e.g., presenting an /ao/ syllable and comparing the pattern of results).

These results thus agree with the abovementioned notion that, due to the low-pass filter characteristics of the womb, fetuses are probably isolated from the mid and high frequency acoustic content of external sounds that characterizes most of the temporal fine structure of speech, \textsuperscript{46,70}. Yet, while lacking the required prior experience for a mature perceptual system responding accurately to high frequencies, the ability to encode fine structure per se seems to be present at birth. Future testing with premature babies early exposed to natural sounds may shed more light on this issue.

Overall, our results are in line with the idea that humans, despite their limited experience to speech at birth, present mature functional mechanisms to detect changes in speech features at an unexpectedly early age,\textsuperscript{4,44}, and since alterations in the neural mechanisms underlying temporal envelope encoding are associated to several disabilities such as autism,\textsuperscript{48} dyslexia,\textsuperscript{49} or other learning problems,\textsuperscript{53}, it is tempting to speculate that the encoding of temporal envelope information, such as its periodicity, may play a crucial role in the very first stages of language acquisition.\textsuperscript{50} Temporal envelopes could provide a neural synchrony channel onto which separate neural representations of other speech features would anchor as parts of an ensemble that would, ultimately, give rise to a coherent unitary entity.\textsuperscript{85} Furthermore, there is increasing evidence that the FFR is a brain response that receives subcortical and cortical contributions in a frequency-specific manner, with frequencies below 150 Hz originating mainly from subcortical sources.\textsuperscript{30,82,83,86,87} Therefore, it is tempting to speculate that the effects observed here may reflect the increasing maturation of the subcortical auditory system from birth to adulthood.

The reported differences in formant structure encoding abilities found between newborns and adults open a window of opportunity to study the developmental progression of these skills. Considering that the gradual increase of phase-locking to high-frequencies is age-dependent,\textsuperscript{42} understanding how inter-individual differences in development as revealed by FFR\textsubscript{ENV} neural responses relate to the acquisition of formant encoding perceptual skills could be used to identify potential risks of future disabilities. Early impairment detection is thus critical to allow early interventions and to maximize the development of speech and listening competences, essential requirements for the acquisition of optimal literacy skills.\textsuperscript{45}

### Considerations on speech stimuli commonly used for newborn FFR studies.

In language FFR studies, the most commonly applied speech stimuli are mandarin syllables following the four different lexical tones,\textsuperscript{8,13,48,51,52,88,89} and different single vowels with rising pitch.\textsuperscript{9,10,43,90} The use of these stimuli focused the research field on assessing voice pitch encoding, putting the assessment of formant structure encoding aside. A notable exception is the widely used consonant–vowel syllable /da/\textsuperscript{11,14,42,44,45,49,56,91,92}, which contains a fine structure change during the consonant–vowel transition. The relevance of using this stimulus relies on the fact that stop consonants are an important constraint in populations with literacy impairments,\textsuperscript{53} and since stop bursts are rapid and low in amplitude in the /d/ consonant compared to vowels, even normal-hearing adults and children can find difficult to discriminate it from other contrastive stop consonants.\textsuperscript{32} However, the short duration of the consonant transition and the high (and changing) frequency peak of the formants that compose it (e.g., the difference between /d/ and /g/ appears in the second formant: /da/ F\textsubscript{2} = 1438–1214 Hz, /ga/ F\textsubscript{2} = 1801–1214 Hz), render this type of stimuli suboptimal in the characterization of FFR responses, which exhibit a spectral power decay with increasing frequency,\textsuperscript{82} especially in populations with an immature encoding of the high frequency content of sounds, such as newborns.\textsuperscript{9,42} Hence, while the phase locking to lower frequency sounds could in principle be safely assessed from the first hours of life,\textsuperscript{42,50} as we demonstrate here as well, the lack of prenatal experience to the high frequency content of sounds and the requirement of a later and greater maturation of the auditory system to encode them,\textsuperscript{39,42,46,50} pose some limitations in the design of stimuli suited to study formant structure encoding.

Therefore, we believe our newly designed /oa/ stimulus, with pitch variation and two vowel sections with different formant structure based on relatively lower frequency harmonic components and suitable durations for accurate spectral analyses, enables a proper assessment of speech sound temporal envelope (FFR\textsubscript{ENV}) and temporal fine structure (FFR\textsubscript{FFRTFS}) encoding.

### Conclusion

The present study provides the first evidence that neonates are able to encode not only the voice pitch of speech sounds and its changes with great accuracy, as has been demonstrated in previous research, but also the formant structure. Specifically, newborns show emerging formant structure encoding skills at lower frequency ranges but still immature encoding precision at higher frequency ranges. In addition, having already proved the feasibility of successfully recording temporal envelope and temporal fine structure in newborns, we here promote the use of this new stimulus as a powerful tool to perform a longitudinal assessment of speech encoding in babies from their very first hours of life throughout the first years of infant development.

### Methods

#### Participants.

A sample of 34 healthy term newborns (17 females; mean gestational age = 40.19 ± 1.08 weeks; mean birth weight = 3379 ± 289 g; aged 14–78 h after birth) was recruited from Sant Joan de Déu Hospital in Barcelona (Spain). Obstetric pathologies, high-risk gestations and risk factors related to hearing impairments...
quency range than those characteristic of cortical sources (beyond 100 Hz), and that attentional modulations of the analyzed frequency content of neural responses recorded in the present study belongs to a higher fre-
duration of rejected sweeps), plus recording preparation time (around 5 min). Adult participants were tested when the newborn was asleep again. The total mean duration of a test session was approximately 25 min (two
ernences in stimulus intensities were chosen for the same reason as in click stimulus.
Procedure. All newborns were recorded at the hospital room where they were resting with their mother. After the neonate passed the universal hearing screening test, the researcher started the recording session as soon as the newborn fell asleep, interrupting it to any sign of discomfort or sleep disruption and resuming it when the newborn was asleep again. The total mean duration of a test session was approximately 25 min (two

Data acquisition. FFRs were recorded from both newborns and adults with a SmartEP platform including the cABR and Advanced Hearing Research modules connected to a Duet amplifier (Intelligent Hearing Systems, Miami, Fl, USA), using three disposable snap Ag/AgCl electrodes placed in a vertical montage (ground electrode at the forehead; active at Fpz; online reference at the right mastoid, ipsilateral to the stimulated ear). All electrode impedances were kept < 7 kΩ. The continuous signal was acquired at a sampling rate of 13,333 Hz with an online bandpass filter from 30 to 1500 Hz and epoched from −40.95 (pre-stimulus period) to 249.975 ms relative to stimulus onset. A total of 4000 artifact-free responses were obtained for each participant after automatic rejec-
tion of any sweep with voltage values exceeding ± 30 µV.

FFR processing. Data was bandpass filtered offline from 80 to 1500 Hz. In order to assess voice pitch encod-
ing, it was necessary to accentuate the FFR components corresponding to the encoding of the stimulus envelope, such as the fundamental frequency (F0). Thus, neural responses were averaged by adding sweeps corresponding to the two stimulus polarities [(Rarefaction + Condensation)/2], yielding the envelope-following response

Stimulus. Inspired by the aforementioned previous stimuli limitations (e.g., short duration of consonant transitions and changing formants, high frequency content), a 250 ms two-vowel syllable stimulus with a rising pitch ending (/oa/) was created in Praat (Fig. 1a). The /o/ vowel section (F1 = 452 Hz; F2 = 791 Hz) lasted from 0 to 80 ms, the /a/ vowel section (F1 = 678 Hz; F2 = 1017 Hz) from 90 to 250 ms, and the /oa/ formant transition section from 80 to 90 ms. Stimulus pitch was kept steady at F0 = 113 Hz from 0 to 160 ms and increased linearly up to 154 Hz from 160 to 250 ms. We used 113 Hz F0 instead of the common 100 Hz F0 to avoid electric line noise harmonics by the European 50 Hz alternating current. In order to maximize the detection of differences in vowel formant encoding in the FFR, formant peak frequencies coincided with harmonics of the fundamental. Stimuli were delivered monaurally to the right ear with a stimulus-onset asynchrony (SOA) of 295 ms, in alternating polarities, at an intensity of 65 dB SPL for adults (Etymotic shielded earphones of 300 Ω, ER, Elk Grove Village, IL, USA) and 60 dB SPL for newborns (same earphones connected to a Flexicoupler disposable adaptor, Natus Medical Incorporated, San Carlos, CA) using Intelligent Hearing Systems (Miami, Fl, USA). Differences in stimulus intensities were chosen for the same reason as in click stimulus.

Procedure. All newborns were recorded at the hospital room where they were resting with their mother. After the neonate passed the universal hearing screening test, the researcher started the recording session as soon as the newborn fell asleep, interrupting it to any sign of discomfort or sleep disruption and resuming it when the newborn was asleep again. The total mean duration of a test session was approximately 25 min (two

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group differences (i.e., whether newborns showed different spectral amplitudes of the signal at F0 peak than adults). Since the obtained values were normally distributed, we employed a two-samples T-test to assess for significant differences (10–160 ms), due to the continuous variation in pitch frequency throughout the rising section (160–250 ms).

Spectral amplitude at F0 peak. Spectral amplitude at F0 peak (113 Hz) was calculated as an indicator of the magnitude of neural phase-locking at that specific frequency only during the steady pitch section of the stimulus (10–250 ms), selecting the time lag that corresponds to the maximum cross-correlation value. The obtained values were non-normally distributed, so a Mann–Whitney U test was used to assess for significant group differences (i.e., whether newborns showed a different transmission delay than adults).

Neural transmission delay. Neural lag. Neural lag was taken as an estimation of FFR latency due to the auditory system's neural transmission delay, and was extracted from a cross-correlation of the entire stimulus with the neural response (10–250 ms), selecting the time lag that corresponds to the maximum cross-correlation value. The obtained values were non-normally distributed, so a Mann–Whitney U test was used to assess for significant group differences (i.e., whether newborns showed a different transmission delay than adults).

Voice pitch encoding. To determine the abilities of newborns (by comparison with adults) to encode the voice pitch contour of the auditory stimulus presented, several parameters were extracted from the FFRcENV:

- **Spectral amplitude at F0 peak.** Spectral amplitude at F0 peak (113 Hz) was calculated as an indicator of the magnitude of neural phase-locking at that specific frequency only during the steady pitch section of the stimulus (10–160 ms), due to the continuous variation in pitch frequency throughout the rising section (160–250 ms). Since the obtained values were non-normally distributed, we employed a two-samples T-test to assess for significant group differences (i.e., whether newborns showed different spectral amplitudes at the F0 peak than adults).

- **Signal-to-noise ratio.** Signal-to-noise ratio (SNR) at F0 peak was taken as an estimation of the relative spectral magnitude of the response, taking into account not only the amplitude value of the signal at the frequency peak of interest (113 Hz) but also around that peak. Therefore, we divided the mean amplitude within a ± 5 Hz frequency window centered at the peak of the frequency of interest (F0) by the mean amplitude within two 28 Hz wide frequency windows (flanks) centered at ± 19 Hz from the frequency of interest (e.g., for F0 = 113 Hz, the mean amplitude from 108 to 118 Hz divided by the average of the mean amplitude from 80 to 108 Hz and the mean amplitude from 118 to 146 Hz). In order to ascertain group differences in the magnitude of the FFR encoding and discern whether newborns had different responses to voice pitch than adults, we used Mann–Whitney U tests because the obtained values were non-normally distributed.

- **Stimulus-to-response cross-correlation.** In order to assess the accuracy with which the FFRcENV reproduces the stimulus waveform, we calculated the normalized cross-correlation between each individual's neural response and the stimulus, separately for the /a/ steady (90–160 ms) and /a/ rising pitch contour stimulus sections (160–250 ms). The maximum value reached within a time lag of 3 to 10 ms (corresponding to the neural lag) was selected (Pearson’s r; values from −1 to 1). The obtained values were non-normally distributed. Therefore, to test for putative between-subjects differences (i.e., whether newborns showed different overall stimulus–response correlation than adults), a Mann–Whitney U test was used, with Age (newborns; adults) as grouping variable and Stimulus Section (/a/ steady; /a/ rising) as contrast variable. To test for putative within-subjects differences (i.e., whether stimulus–response correlations were different depending on stimulus pitch contour), a Wilcoxon test for two related samples comparing the correlation values obtained for each stimulus section (/a/ steady; /a/ rising) was used. Finally, to test for a putative interaction between factors (i.e., whether newborns showed a different correlation value depending on stimulus pitch section than adults), a Mann–Whitney U test was used taking Age (newborns; adults) as grouping variable and the difference between the two conditions of the Stimulus Section (/a/ steady – /a/ rising) as contrast variable.

- **Pitch error.** Pitch error per stimulus section was used to determine pitch-tracking accuracy of the F0 contour (corresponding to the autocorrelation peak lag per bin) by averaging the absolute Euclidian distance between the stimulus F0 contour and the response F0 per pitch section separately (steady [10–160 ms]; rising [160–250 ms]; starting from the onset of the section + the individual neural lag; values in Hz). Since obtained values were non-
normally distributed, to determine between-subject effects, within-subjects effects and interaction, we followed the same procedure as with the stimulus-to-response cross-correlation explained above.

**Pitch strength.** Pitch strength per stimulus section was taken as a measure of periodicity and the magnitude of neural phase-locking of the response\(^{10}\), and calculated by averaging the obtained peak autocorrelation value of the response across bins, per pitch section separately (steady; rising; starting from the onset of the section + the individual neural lag; values from ~ 1 to 1). Values were non-normally distributed, thus an identical method with the same factors as employed above in cross-correlation and pitch error parameters was used to determine between-subject effects, within-subjects effects and interaction.

**Formant structure encoding.** Regarding the encoding of the perceptual quality of formant structure, several parameters were retrieved from the FFR\(_{250}\).

**Spectral amplitude.** Spectral amplitudes at spectral peaks corresponding to stimulus F\(_1\) frequencies (452 Hz [\(/o/\)] and 678 Hz [\(/a/\)]) were retrieved separately from neural responses to the \(/o/\) section (10–80 ms) and the \(/a/\) steady section (90–160 ms). All values were normally distributed, so an ANOVA test was conducted. Regarding the spectral amplitude at 452 Hz, (a) the Group variable (newborns; adults) was chosen as between-subjects factor, to examine whether newborns showed different amplitude values at 452 Hz than adults; (b) Stimulus Section (\(/o/\) section; \(/a/\) section) as within-subjects factor, in order to test whether spectral amplitudes at 452 Hz were different depending on stimulus vowel section; (c) Interaction between factors was analyzed to ascertain whether newborns showed a different amplitude value at 452 Hz depending on stimulus vowel section than adults. Pursuing an identical purpose, we conducted again the same test to examine differences at 678 Hz. The transition from \(/o/\) vowel to \(/a/\) vowel was not analyzed due to its short duration (10 ms).

**Signal-to-noise ratio.** Following the same procedure as with the spectral amplitude, SNRs at spectral peaks corresponding to stimulus F\(_1\) frequencies (452 Hz [\(/o/\)] and 678 Hz [\(/a/\)]) were also retrieved separately from responses to the \(/o/\) and the \(/a/\) steady section, using an identical method to calculate it as described above for the FFR\(_{250}\). All values were normally distributed, so ANOVA tests on 452 Hz and 678 Hz were conducted with the same factors and objectives as described above for F\(_1\) spectral amplitudes analyses.

All analyses were additionally computed by excluding participants with extreme values (more than three interquartile ranges; \(N = 9; 4\) newborns + 5 adults). As the statistical results obtained did not alter the main findings of the study, we decided to keep all participants within the reported analyses to better represent the inherent variability of our samples (results excluding extreme values are reported in Suppl. Tables 2–7).

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**Competing interests**
The authors declare no competing interests.

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