

# Development of a small wireless device for perspiration monitoring

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1 **Development of a small wireless device for perspiration monitoring**

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18 **Abstract**

19 A small and wireless device that can capture the temporal pattern of perspiration by a novel  
20 structure of water vapor collection combined with reusable desiccant has been developed. The  
21 novel device consists of a small cylindrical case with a temperature/relative humidity sensor,  
22 battery-driven data logger, and silica gel (desiccant). Water vapor of perspiration was detected  
23 by the change in relative humidity and then adsorbed by silica gel, allowing continuous  
24 recording of perspiration within a closed and wireless chamber, which has not been previously  
25 achieved. By comparative experiments using the commercially-available perspiration  
26 monitoring device, the developed device could measure perspiration as efficiently as the  
27 conventional one, with a normalized cross coefficient of 0.738 with 6 s delay and the interclass  
28 correlation coefficient [ICC(2, 1)] of 0.84. These results imply a good agreement between the  
29 conventional and developed devices, and thus suggest the applicability of the developed device  
30 for perspiration monitoring.

31

32 **Key words:** perspiration monitoring, emotional sweating, sympathetic activity

33

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38 **1. Introduction**

39 Perspiration, or sweating, is one of the most fundamental phenomena in human physiological  
40 events. The main result of systemic sweating is cooling effect for thermoregulation by sweat  
41 evaporation [1], which is called “thermal sweating”. There is another type of sweating called  
42 “emotional (or mental) sweating.” Emotional stresses (e.g., rising tension, upset, and  
43 concentration) trigger sweating, particularly on the face, palm, and sole via sympathetic nervous  
44 tone [2-4]. To date, a number of diseases have been reported to be associated with sweating  
45 abnormalities such as thyroid diseases [5], dysautonomia [6], menopause [7], and social anxiety  
46 disorder [8]. In addition, the perspiration monitoring can be utilized for prediction or diagnosis  
47 of nervous disorders such as brachial plexus avulsion (BPA) [9] and reflex sympathetic  
48 dystrophy [10]. Especially, the monitoring of sympathetic activity such as perspiration might be  
49 important for the early diagnosis of obstetric BPA [11], because it is often difficult for neonates  
50 to express their symptoms verbally. In light of the possible applications of ubiquitous  
51 perspiration monitoring such as a prediction or diagnosis of perspiration-related disorders, a  
52 small, convenient, and sensitive device for perspiration monitoring has been desired.

53           A skin conductance meter was used as an indirect method for estimation of sweating  
54 [12], and a simple humidity meter was employed to measure water evaporation from the skin  
55 [13,14]. Recently, wearable, adhesive, and tattoo-like sweat monitoring device have been  
56 proposed [15-17], although they are more intended for prediction of sweat electrolytes rather  
57 than perspiration amount, or they give an indirect index of perspiration. At this time the latest  
58 perspiration measurement device involves colorimetric detection by a digital camera, which  
59 requires special setup [18]. As a more direct measurement of water exchange, the vapor pressure  
60 diffusion method and ventilated chamber method were developed [19,20]. The vapor pressure  
61 method utilizes the theory that the amount of water exchange ( $F$ ) in natural flow is calculated as  
62  $F = D(\partial p/\partial x)$ , where  $D$  is the temperature- and atmospheric pressure-dependent diffusion  
63 coefficient,  $p$  is the water vapor pressure, and  $x$  is distance from the surface [19]. Because  $p$  can

64 be calculated from relative humidity and temperature, at least four sensors [(humidity +  
65 temperature) × two points] are required. In addition, this method relies on the assumption that  
66 the state of the outer atmosphere is unchanging, which is unlikely in daily perspiration  
67 monitoring. To address this, a closed chamber system with water vapor condenser was proposed  
68 [21,22], although the coolant (Peltier device) is required and thus power consumption would be  
69 measurable. The theory of the ventilated chamber method is similar to that of the vapor pressure  
70 method, except this method uses forced and constant airflow. The constant airflow is injected  
71 into a small chamber adjacent to the skin, and the air with evaporated water vapor is collected in  
72 an outlet chamber. The amount of water exchange is then calculated with the airflow rate and  
73 difference of humidity between inlet and outlet air [19,20,23]. However, it is difficult to contain  
74 air ventilator and chambers in one small package such that it can be of practical use in daily life.  
75 Therefore, it was considered beneficial to develop a small device that could monitor  
76 perspiration and allow prediction of emotional and physiological status.

77           The aim of this study was to develop a small device for perspiration monitoring and  
78 compare its performance with a conventional sweat meter and stress analyzing method under  
79 conditions of mental stress.

80

## 81 **2. Materials and Methods**

### 82 *2.1. Developed device*

83 Fig. 1 shows the exterior (Fig. 1A, B) and structure (Fig. 1C) of the developed device. A  
84 custom-made data logger circuit with battery, dry silica-gel (Wako Pure Chemical Industries,  
85 Ltd., Osaka, Japan), and a small temperature/relative humidity (T/RH) sensor (SHT-21,  
86 Sensirion AG, Zürich, Switzerland; accuracy of temperature is  $\pm 0.3^{\circ}\text{C}$ , accuracy of relative  
87 humidity is  $\pm 2\%$ , calibrated at Industrial Research Institute of Ishikawa, Japan) with a sampling  
88 rate of 1 Hz was encapsulated in this order in a small plastic chamber toward measuring  
89 windows facing the skin (Fig. 1C).

90

## 91 2.2. Perspiration rate calculation

92 The theory of this equipment is based on the vapor pressure method [19,21] with modifications.

93 Fig. 2 illustrates the method of perspiration measurement in the developed equipment.

94 According to the Fick's law of diffusion, the flux of water vapor  $J$  ( $\text{g m}^{-2} \text{s}^{-1}$ ) between two  
95 points can be calculated as Eq. (1):

$$J = -D \frac{dH}{dx} \quad (1)$$

96 where  $D$  ( $\text{m}^2 \text{s}^{-1}$ ) is a diffusion coefficient of water vapor in the air,  $dH$  ( $\text{g m}^{-3}$ ) is a difference of  
97 concentration of water vapor, and  $dx$  (m) is a distance between two points. In the developed  
98 equipment, two different fluxes of water vapor: from the skin surface to the T/RH sensor (Fig.  
99 2A, green arrow; **s-w**), and from the sensor to the dry chamber (Fig. 2A, blue arrow; **w-d**), can  
100 be theorized. The flux difference between (**s-w**) and (**w-d**) could be detected as a change of  
101 humidity in T/RH sensor. The water exchange between the skin and silica gel via T/RH sensor  
102 should satisfy Eq. (2):

$$\begin{aligned} V \frac{\Delta H_x(t)}{\Delta t} &= A_1 J_1 - A_2 J_2 \\ &= A_1 D \frac{H_1(t) - H_x(t)}{L_1} - A_2 D \frac{H_x(t) - H_2}{L_2} \end{aligned} \quad (2)$$

103 where  $V$  ( $\text{m}^3$ ) is a volume of wet chamber in which the T/RH sensor exists;  $H_1(t)$ ,  $H_x(t)$ , and  $H_2$   
104 ( $\text{g m}^{-3}$ ) are the concentrations (i.e., absolute humidity) of water vapor at the skin surface, T/RH  
105 sensor, and dry chamber, respectively;  $A_1$  and  $A_2$  ( $\text{m}^2$ ) are the areas of windows at (**s-w**) and  
106 (**w-d**) junctions, respectively;  $J_1$  and  $J_2$  are the fluxes of (**s-w**) and (**w-d**), respectively;  $L_1$  and  
107  $L_2$  (m) are the distances of (**s-w**) and (**w-d**), respectively (Fig. 2A).  $H_2$  is assumed to be  
108 constant due to a buffering effect of desiccant (preliminary experiment is shown in Fig. S1).  
109 Because  $J_1$  in Eq. (2) simply represents the flux of total water vapor from the skin surface [i.e.,  
110 perspiration and constant transepidermal water loss (TEWL)], the Eq. (2) can be solved for  $J_1$  as  
111 following Eq. (3):

$$W(t) = J_1 = \frac{V}{A_1} \frac{\Delta H_x(t)}{\Delta t} + \frac{A_2 D(t)}{A_1 L_2} (H_x(t) - H_2) \quad (3)$$

112 where the rate of water vapor diffusion from the skin  $W(t)$  ( $\text{g m}^{-2} \text{s}^{-1}$ ) can be calculated only by  
 113 measuring  $H_x(t)$  with T/RH sensor, as  $V$ ,  $A_1$ ,  $A_2$ ,  $L_2$ , and  $H_2$  are all considered fixed, and  $D(t)$  can  
 114 be calculated by using following Eq. (4) under normal atmospheric pressure [24]:

$$D(t) = 1.87 \times 10^{-10} \times T(t)^{2.072} \quad (4)$$

115 where  $T(t)$  (K) is the temperature at the time  $t$ .

116 Here, the fixed values were set to:  $V = 6.3 \times 10^{-7} \text{ m}^3$ ,  $A_1 = 1.3 \times 10^{-5} \text{ m}^2$ ,  $A_2 = 5.6 \times 10^{-5} \text{ m}^2$ ,  $L_2$   
 117  $= 5.0 \times 10^{-3} \text{ m}$  according to the equipment design, and  $H_2$  was estimated to be  $1.7 \text{ g m}^{-3}$  (Fig.  
 118 S1). Finally, the perspiration  $Per(t)$  ( $\text{mg cm}^{-2} \text{ min}^{-1}$ ) [20,23] can be calculated from  $W(t)$  by a  
 119 simple conversion Eq. (5):

$$Per(t) = 6W(t) \quad (5)$$

120 because  $1 \text{ g m}^{-2} \text{ s}^{-1}$  is equal to  $6 \text{ mg cm}^{-2} \text{ min}^{-1}$ .

121 The conversion from relative humidity  $h$  (%) to absolute humidity  $H$  ( $\text{g m}^{-3}$ ) was as  
 122 following Eq. (6) based on the ideal gas law:

$$H = \frac{M_w P_s(T)}{RT} \frac{h}{100} \quad (6)$$

123 where  $M_w$  is the molecular weight of water ( $= 18.02 \text{ g mol}^{-1}$ ),  $P_s(t)$  is the saturated water vapor  
 124 pressure (kPa) at the temperature  $T$  (K), and  $R$  is the gas constant ( $R = 8.314 \times 10^{-3} \text{ kPa m}^3 \text{ K}^{-1}$   
 125  $\text{mol}^{-1}$ ), according to the American Society of Heating, Refrigerating and Air-Conditioning  
 126 Engineers guidelines [25].

127 Because the change in water vapor flux includes constant water loss [26] and  
 128 perspiration, a baseline subtraction has been employed. As shown in Fig. 2B, it is theorized that  
 129 the baseline (i.e., lower envelope) and wave crests reflect the constant water loss and  
 130 perspiration, respectively. To extract the perspiration pattern after the recording of water vapor  
 131 flux, the difference between water vapor flux and the baseline was calculated by the embedded  
 132 program in Origin software (version 2015; OriginLab Corp., MA, USA).

133

134 *2.3. Verification of developed equipment*

135 The performance test, in which the direct water vapor was applied to the developed device, was  
136 first performed (for details see Fig. S2). For verification of the developed perspiration monitor  
137 and calculation method described above, the perspiration pattern obtained from the developed  
138 device and commercially available conventional sweat meter was compared as follows. First,  
139 five individuals for the test were employed after obtaining written informed consent. The two  
140 devices, both newly developed and conventional devices (SKD-1000, Skinoss Co., Ltd., Nagano,  
141 Japan; nominal uncertainty is  $\pm 10\%$  of measured value), were attached side-by-side to the palm  
142 of each individual, followed by simultaneous perspiration recording in a sitting position for 30  
143 min. After recording, the temporal changes of perspiration were compared using a normalized  
144 cross-correlation function (nCCF). The corresponding peak-to-peak values of perspiration  
145 patterns were analyzed by general Deming regression [27], absolute interclass correlation  
146 coefficient of two variables [ICC(2, 1)] [28,29], and Bland–Altman plot [30,31] to estimate the  
147 agreement of both devices.

148

149 *2.4. Measurement of perspiration evoked by sympathetic activity*

150 To determine if the developed equipment could detect perspiration under stress conditions, an  
151 experiment was conducted utilizing sympathetic activity. First, the developed device and skin  
152 potential sensor (NE-114A; Nihon Kohden Corp., Tokyo, Japan) were attached to their palm  
153 (Fig. 4A). Furthermore, they were requested to perform the following tasks: (1) take a deep  
154 inspiration 5 times at intervals of 1 min and (2) do a mental calculation (e.g., the subjects were  
155 orally requested to continuously subtract 7 from 100) for 5 min to evoke sympathetic activity  
156 that involves perspiration on the palm [32,33]. During the test, the perspiration and sympathetic  
157 skin response (SSR) on the palm were recorded [33]. These data were used to confirm if the  
158 developed device could detect perspiration by mental stress.

159

160 2.5. *Ethical approval*

161 These protocols involving human subjects were approved by the Medical Ethics Committee of  
162 Kanazawa University (#553).

163

164 **3. Results**

165 3.1. *Perspiration recording by the developed and conventional devices*

166 As a result of performance test, the developed device was confirmed to be able to measure the  
167 water exchange with an uncertainty of  $<\pm 5\%$  and a long-term stability of  $>4$  h (Fig. S2). The  
168 representative temporal changes in perspiration recorded by the developed and conventional  
169 devices are shown in Fig. 3A, B (the baseline data were shown in Fig. S3). The comparable time  
170 profiles of perspiration in a steady state were observed (Fig. 3A, B). Perspiration pattern  
171 determined by the developed device was in good agreement with that of the conventional device  
172 [Fig. 3C; peak correlation coefficient of 0.738 at  $-6$  s, and Fig. 3D; ICC(2, 1) of the  
173 corresponding peak-to-peak amplitudes was 0.84 with the 95% confidence interval (CI) of  
174 0.76–0.90]. The Bland–Altman plot revealed a fair agreement between both devices [Fig. 3E;  
175 bias =  $-0.0042$  mg cm<sup>-2</sup> min<sup>-1</sup> (95% CI:  $-0.014$ – $0.0056$ ) with the limits of agreement  $-0.087$  to  
176  $0.079$ ]. These results imply that the developed device could capture perspiration as efficiently as  
177 the conventional one. The other subjects showed similar results (data not shown).

178

179 3.2. *Detection of palmar perspiration evoked by the sympathetic activity using the developed*  
180 *device*

181 Furthermore, whether the developed device could capture the onset of perspiration was tested.  
182 To test this, sympathetic activation that is related to palmar sweating [32,33] was utilized. For  
183 measurement of sympathetic activity, the palmar SSR was recorded at the same time that  
184 perspiration was measured (Fig. 4A). With these sensors, subjects were requested to perform  
185 two tasks (a deep inspiration and mental calculation) to evoke sympathetic activity. During the

186 stress test, the palmar SSR suggested substantial reactions according to the stressor (Fig. 4B,  
187 SSR). In the same manner, palmar perspiration recorded by the developed device showed a  
188 stress-induced pattern with good agreement with the SSR (Fig. 4B, perspiration). From these  
189 results, it is plausible that the developed device can indeed capture the onset of perspiration. The  
190 other subjects showed similar results (data not shown).

191

#### 192 **4. Discussion**

193 The aim of this research was to develop a small device for perspiration monitoring. To achieve  
194 this, a small, stand-alone temperature and relative humidity sensor used to calculate absolute  
195 humidity was designed, allowing wireless monitoring of water exchange with a small exterior.  
196 In addition, a novel closed-chamber system with silica gel allowing constant measurement  
197 independent from the ambient condition was introduced. The developed sensor for perspiration  
198 monitoring was validated in human subjects by a comparison of the conventional and developed  
199 devices and by concurrent monitoring of sympathetic activity-related perspiration.

200 In this study, a modification of the vapor pressure method [19] was utilized. In the  
201 conventional vapor pressure method, water exchange on the skin (i.e., constant water loss and  
202 perspiration) can be detected as the natural flow of water vapor. However, the flow from the  
203 skin to ambient air is dependent on outer air conditions such as temperature and humidity, i.e.,  
204 the previous method would be deeply affected by the nature of outer atmosphere. Thus, a  
205 combination of a closed chamber with enforced ventilation has been developed [20,23,34].  
206 These combined methods use dehumidified nitrogen or ventilation pumps, which could hamper  
207 daily monitoring of perspiration patterns. To address these limitations, a closed-chamber filled  
208 with silica gel above the T/RH sensor was developed (Figs. 1 and 2). In this system, the  
209 adsorption of water vapor into silica gel generates a natural but constant flow of water vapor.  
210 Under such a constant flow, the T/RH sensor below the silica gel can constantly measure the  
211 water exchange without the interference of ambient air in a small and wireless exterior (Figs. 1

212 and 2).

213 In principle, the developed device is relying on the consistency of humidity in a  
214 desiccant-filled chamber [ $H_2$  in Fig. 2A and Eq. (3)]. According to the preliminary study, the  
215 variability of relative humidity in a silica gel-filled chamber was small (Fig. S1; about 1–2 g  
216  $m^{-3}$ ) and can be considered static when compared to the change of humidity in a wet chamber  
217 facing to the skin (about 15–30 g  $m^{-3}$ ). The change of  $H_2$ , therefore, could be negligible. That  
218 said, a dual T/RH sensor system (one sensor is in a wet chamber, the other is in a  
219 desiccant-filled one) would increase an accuracy of perspiration measurement by eliminating  
220 the small fluidity of  $H_2$ , although the power consumption would be doubled and thus,  
221 measurable time would be halved.

222 Water evaporation from the skin includes constant water loss [26] and perspiration.  
223 Among them, perspiration was particularly focused because a number of diseases are associated  
224 with perspiration abnormalities [5-8,35-41]; therefore, monitoring of perspiration would be  
225 considered beneficial for health. Because the water exchange measured by the developed device  
226 includes both constant water loss and perspiration, a baseline subtraction (Fig. 2B) was adopted  
227 to extract perspiration. In this method, it was theorized that the baseline indicates constant water  
228 evaporation, while the “crests” reflect perspiration. As a result of developed methods,  
229 comparable perspiration profiles were obtained between conventional and developed devices  
230 (Fig. 3). The high cross correlation (nCCR = 0.738) and interclass correlation coefficient of the  
231 peak-to-peak values [ICC(2, 1) = 0.84] indicate the considerable agreement between the  
232 conventional and developed equipment (Fig. 3). In addition, the developed device could detect  
233 perspiration evoked by mental stress (Fig. 4) like as the conventional device [32,33], which  
234 further adds to the applicability of this system for perspiration monitoring.

235 It should be noted, however, that there are some limitations with respect to the  
236 developed device.

237 First, the developed device cannot detect constant water loss (e.g., TEWL) that the

238 conventional device can measure (note the baseline shift of perspiration profiles in Fig. 3A).  
239 This is because of the baseline subtraction introduced in this study (Fig. 2B). In principle, the  
240 information about constant water loss including TEWL was eliminated by baseline subtraction.  
241 The data of baseline themselves might have information on TEWL, although further analysis  
242 about the baseline data would be required.

243           Second, the perspiration amount (i.e., the peak-to-peak amplitude of perspiration  
244 profiles) did not always correspond between conventional and developed devices, especially at  
245 higher values (Fig. 3D, E). The slight difference could be explained by the uncertainty of each  
246 variable in Eq. (3) used for perspiration calculation. Although we have determined the overall  
247 variability as being <5% (Fig. S2), the uncertainty of each variable such as  $H_x(t)$  and  $H_2$ , which  
248 has not been estimated in this study, is also the seed of error. A more accurate measurement  
249 could be achieved by incorporating such errors, although the calibration of the T/RH sensor and  
250 the performance test should be enough for general perspiration monitoring. The difference could  
251 also be explained by the responding speed of the T/RH sensor. The sensor used in this study  
252 (SHT-21) has a time constant of 8 s, corresponding to the delay of approximately 8 s in the  
253 output. It is also possible that there is a speed limit of water vapor adsorption in the silica gel. It  
254 is plausible, therefore, that the large and sharp spike of perspiration might not be appropriately  
255 detected by the T/RH sensor because of the sensing delay and/or adsorption speed limit. Indeed,  
256 the perspiration pattern recorded by the developed device reported slightly delayed (6 s) data  
257 compared to the conventional one (Fig. 3C), the waveform of the developed device was blunter  
258 than that of the conventional device, and the response against the decrease of perspiration is  
259 slower than that of the perspiration onset (Fig. 3A, B). Third, the saturation of desiccant would  
260 be a problem. The silica gel used in the device (about 4 g) can capture up to about 1.2 g of water  
261 vapor (i.e., about 30% of its own weight) at body temperature [42]. This capacity could be  
262 sufficient to capture water vapor for >900 minutes if the constant  $10 \text{ mg cm}^{-2} \text{ min}^{-1}$  water vapor  
263 evaporation is simulated, which is beyond the observed perspiration plus water loss (Figs. 3A

264 and S3). In addition, the performance test has proved that at least 4 h continuous water vapor  
265 adsorption did not affect the readout value of the device (Fig. S2). Therefore, the saturation of  
266 desiccant in the developed device would be considered negligible.

267           Despite these drawbacks, the developed device could measure perspiration profiles  
268 in an easy and convenient way, which may be suitable for daily monitoring of perspiration. The  
269 next goal should be to confirm the more precise estimation of perspiration amount for  
270 diagnostic purpose, to analyze the baseline data which may contain the information about  
271 TEWL, and to explore the applicability of the device at various positions on the body;  
272 nonetheless the detection of perspiration at anterior chest has been already confirmed (data not  
273 shown).

274           In conclusion, a small and wireless device was developed to capture the temporal  
275 pattern of perspiration using a novel method of water vapor collection combined with a reusable  
276 desiccant. With further refinement, this system could be applicable for daily perspiration  
277 monitoring, and could predict the onset of the diseases related to perspiration abnormalities.

278

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285 2012-085983.

286

## 287 **Ethical Approval**

288 This study was approved by the Medical Ethics Committee of Kanazawa University (#553-1).

289

290 **Conflict of interest**

291 FM is the president of the Rousette Strategy Inc. where the developed device was assembled.

292 No financial support was received from either FM or the Rousette Strategy Inc.

293

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393

394

395 **Figure legends**

396 **Figure 1** Outline of the developed device. (A, B) Exterior of the device. A small plastic cylinder  
397 (A) contains a temperature/relative humidity (T/RH) sensor, electric boards, and silica gel (B).  
398 (C) Schematic of the device.

399

400 **Figure 2** Principles of perspiration monitoring. (A) In this system, two different water vapor  
401 fluxes were theorized: from the skin surface to the wet chamber (green arrow; **s-w**), and from  
402 the wet chamber to the dry chamber (blue arrow; **w-d**). The perspiration with constant water  
403 loss can be obtained by the calculation of water vapor flux from the skin surface (green arrow).  
404 (B) After obtaining the temporal data of water vapor flux, the baseline subtraction was  
405 introduced to separate perspiration and constant water loss.

406

407 **Figure 3** Comparison of the temporal patterns of perspiration. (A) Perspiration patterns  
408 obtained by the conventional (upper) and developed (lower) devices are shown. (B) These  
409 devices showed similar patterns of perspiration. (C) The normalized cross correlation function  
410 (nCCF) of these devices. Note a good correlation (0.738) between the conventional and  
411 developed devices with a short delay (-6 s). (D, E) Agreement of peak-to-peak values by the  
412 two devices. (D) Scatter plot of conventional (*x*-axis) and developed (*y*-axis) devices with linear  
413 Deming regression (red line) and 95% prediction interval (PI). These devices showed a good  
414 agreement, with a high absolute interclass correlation coefficient [ICC(2, 1) = 0.84]. A blue line  
415 indicates complete agreement. (E) Bland-Altman plot with the limits of agreement (bias =  
416  $-0.0042 \text{ mg cm}^{-2} \text{ min}^{-1}$  with the limits of agreement  $-0.087$  to  $0.079$ ).

417

418 **Figure 4** Detection of sympathetic palmar perspiration. (A) Experimental condition: the subject  
419 was requested to attach the developed device and sensors of sympathetic skin response (SSR)

420 side-by-side followed by maintaining a rest position. (B) Representative data of the SSR and  
421 palmar perspiration. SSR activity was observed by deep inspiration and mental calculation. The  
422 developed device could successfully record palmar perspiration in response to the SSR.

423

424 **Figure S1** The humidity change in wet and dry (desiccant) chambers. Although these data were  
425 separately obtained in the same condition, a very small change of absolute humidity in a dry  
426 chamber (B) compared to the wet one (A) should be noted.

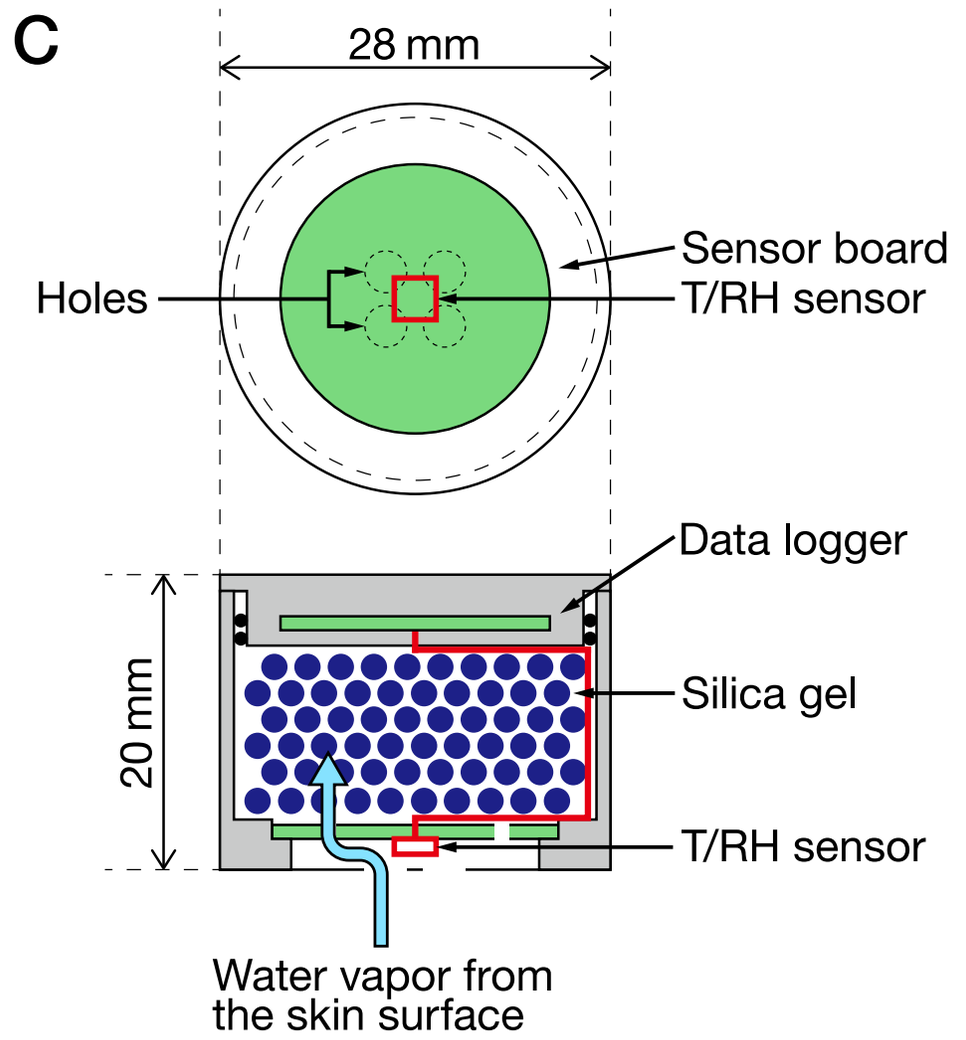
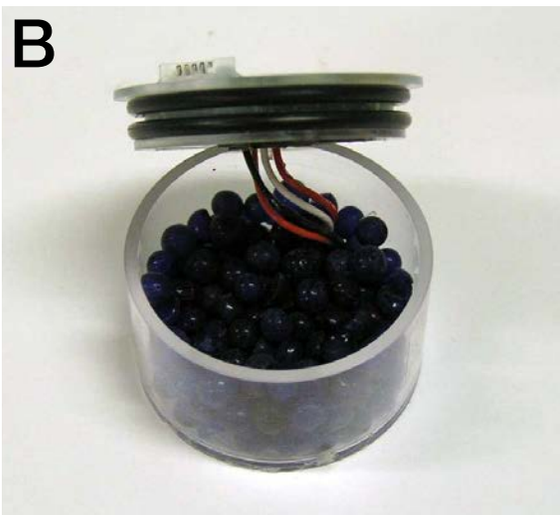
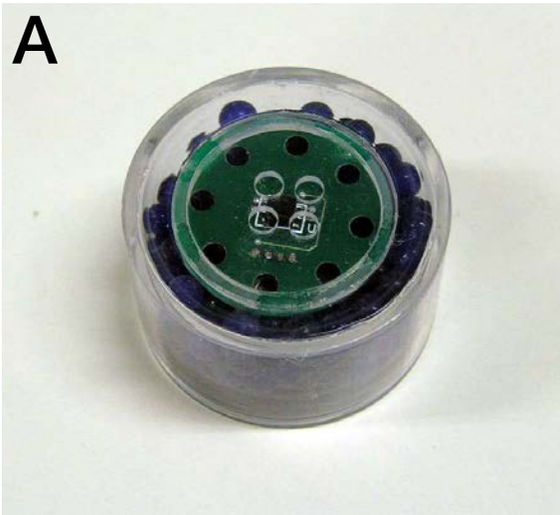
427

428 **Figure S2** The performance test of the developed device. In the test, the developed device was  
429 placed over the cylindrical water tank with a polytetrafluoroethylene (PTFE) sealant (A) in the  
430 temperature-controlled room. The constant and massive water vapor was created by heating the  
431 tank at 35°C, and the recording of water vapor flux was performed. As a result, about 4 h  
432 continuous recordings could be achieved (B). The uncertainty of the calculated water vapor rate  
433 was less than  $\pm 5\%$  for  $\sim 4$  h (average =  $14.89 \text{ g m}^{-2} \text{ min}^{-1}$ , min–max = 14.54–15.55;  
434 corresponding to  $-2.3$ – $+4.4\%$  error against the average). After the experiment, the weight  
435 change of the silica gel (i.e., the amount of water transfer) was 108 mg, whereas the integral of  
436 calculated water vapor flux was 103.47 mg for 4 h; the error was  $-4.2\%$  of the actual value (B).

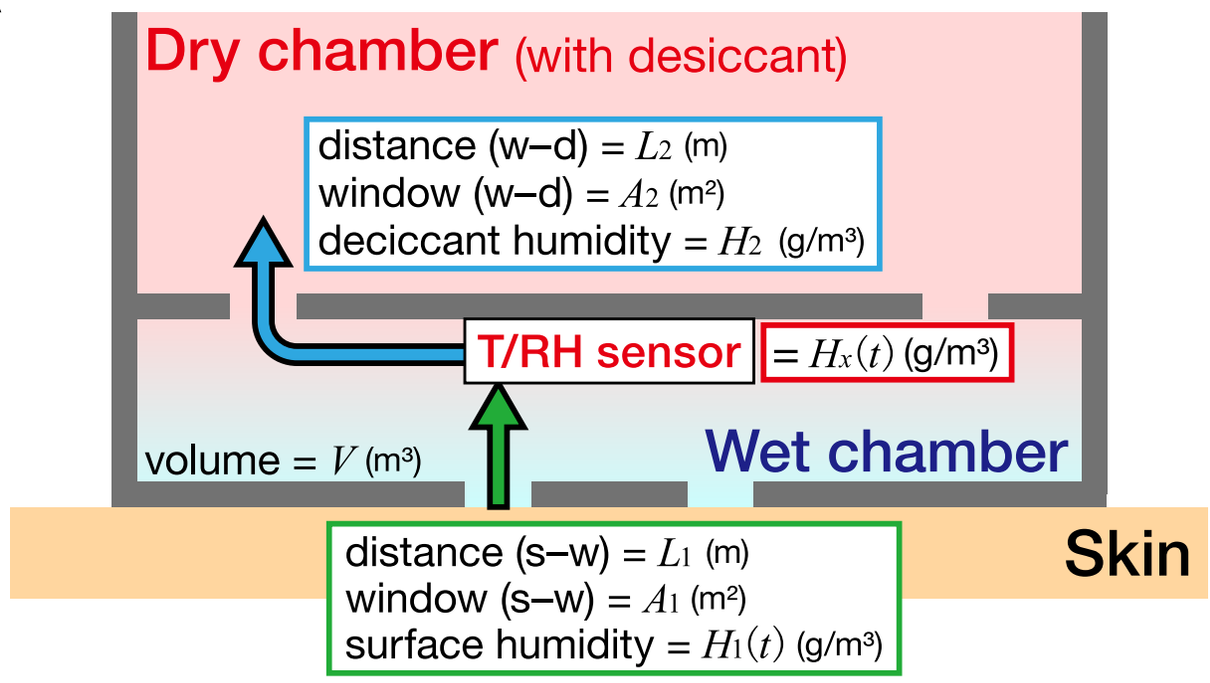
437

438 **Figure S3** The baseline data used in Fig. 3A.

Figure 1



A



B

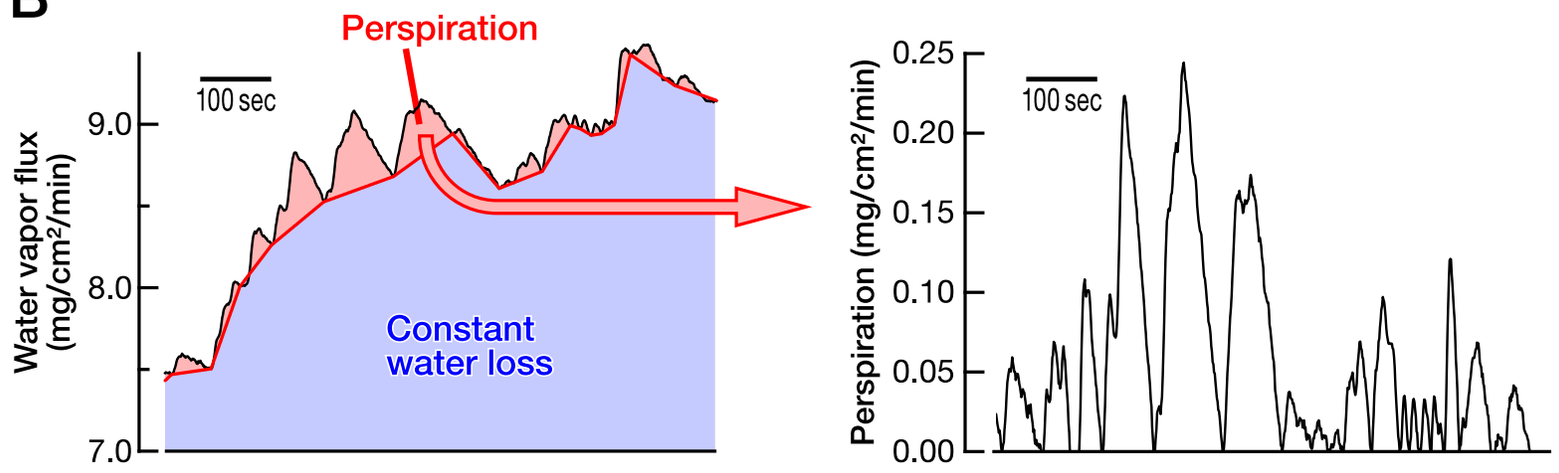
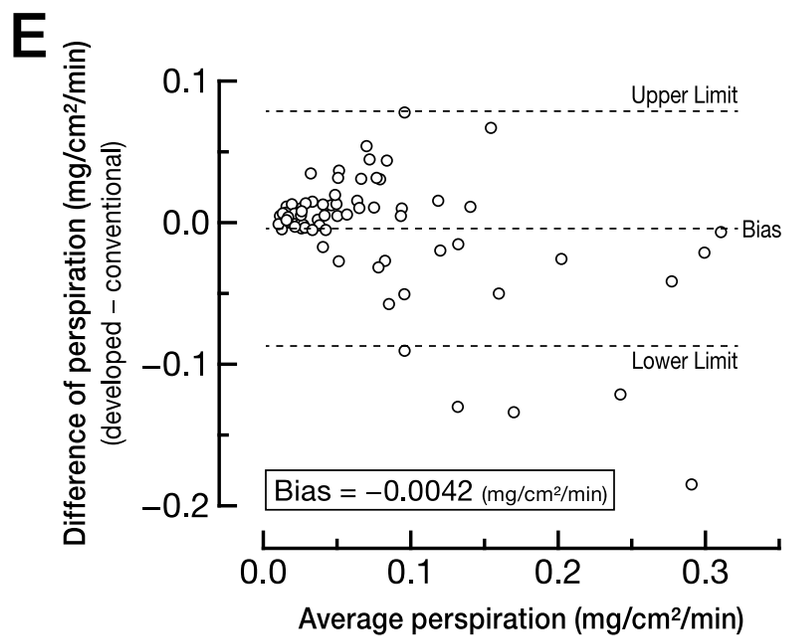
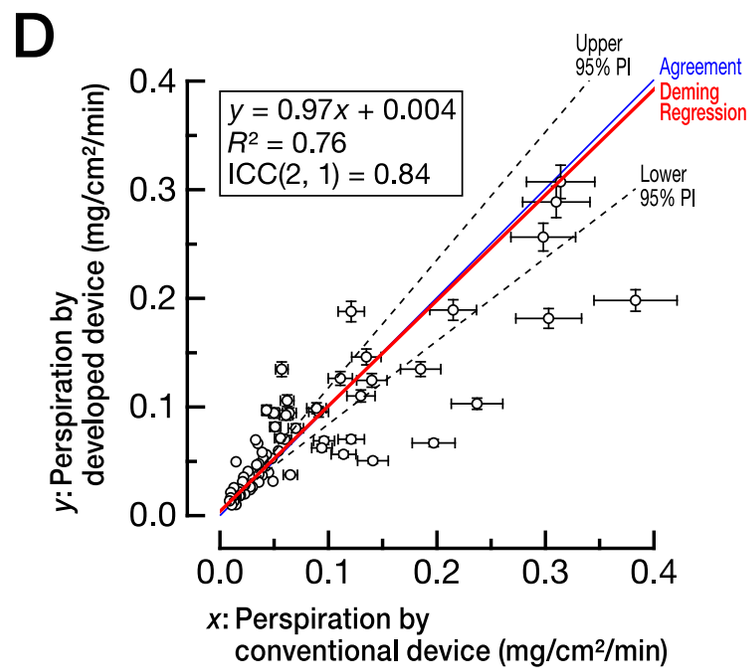
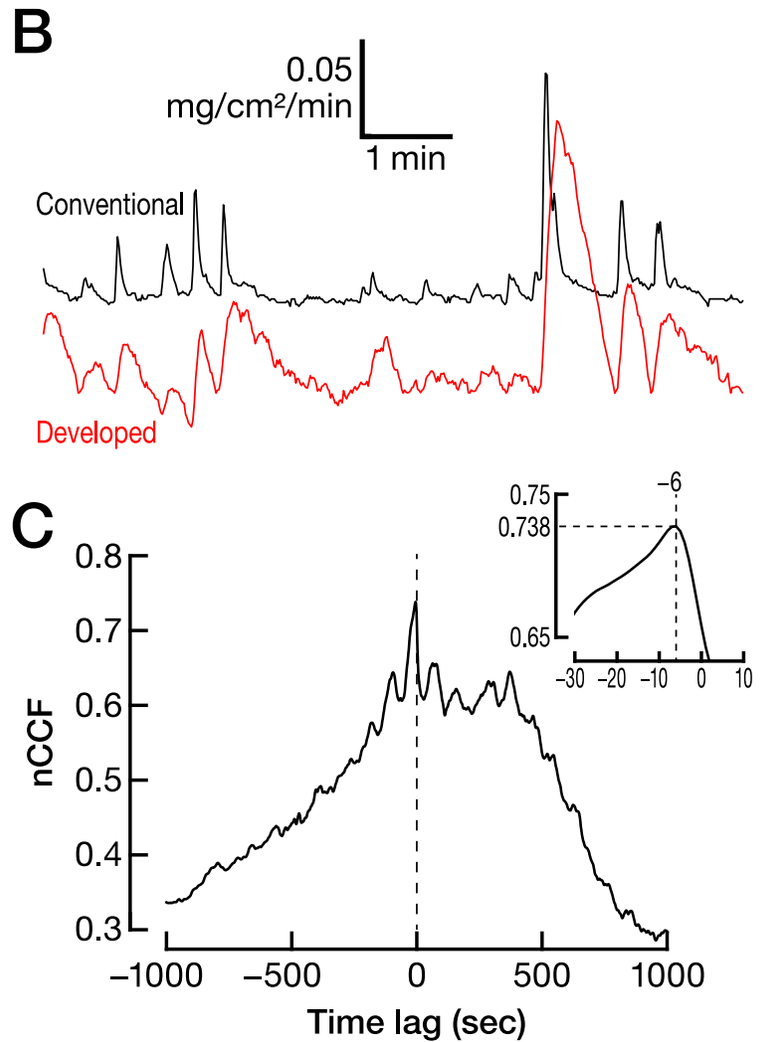
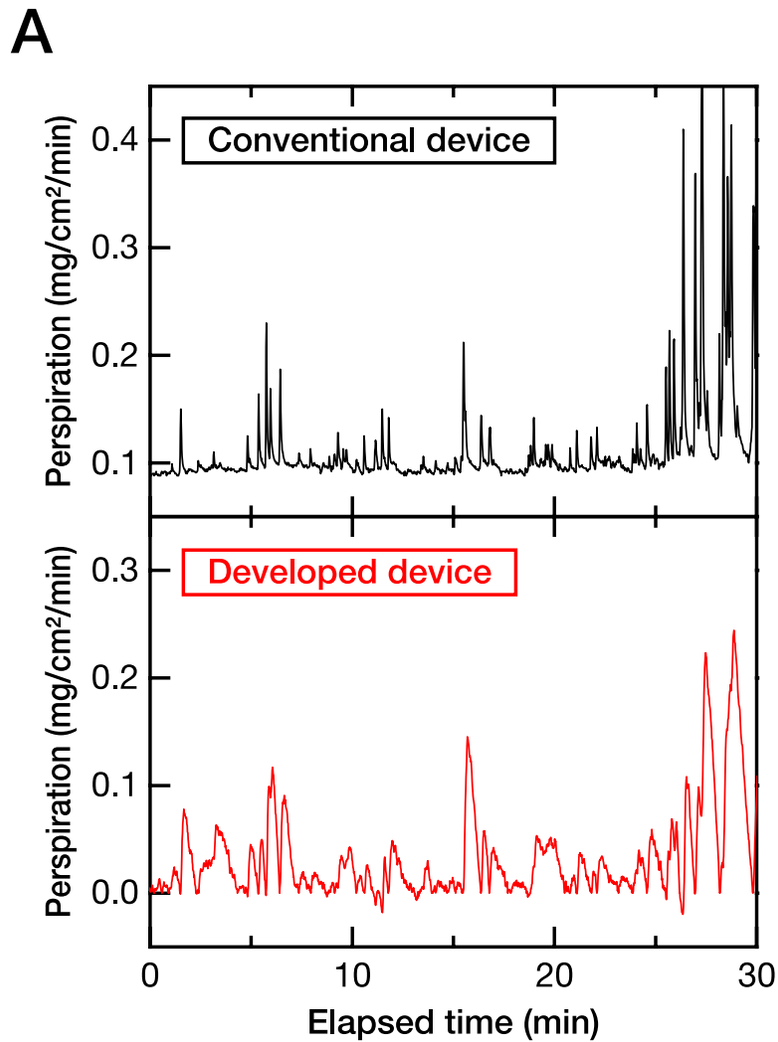


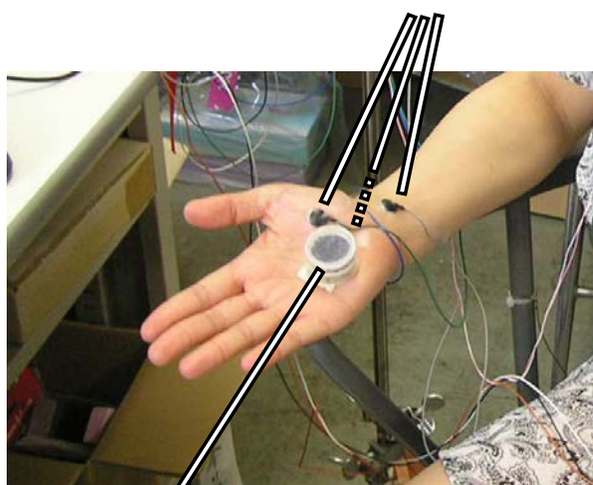
Figure 3



**A**

**SSR sensors**

Two electrodes (on the palm and the back) with an indifferent electrode on the wrist



Developed device

**B**

