MITSUBISHI ELECTRIC RESEARCH LABORATORIES http://www.merl.com

Low-Cost Surface Mount LED Gas Sensor

Roderick L. Shepherd, William S. Yerazunis, King Tong Lau, Dermot Diamond

TR2005-097 October 2005

Abstract

A low-cost chemical sensor comprising surface mount light emitting diodes (LEDs) has been developed for colorimetric based gas detection. The device consists of a pair of LEDs connected to a simple PIC microcontroller circuit, and in most basic form requires the use of only 2 I/O pins on the chip. The key features of this sensor are the use of an LED, rather than a phtodiode for light detection and an all-digital light detection protocol that leads to a reduction in cost and power consumption by avoiding the need for an Analog-to-Digital converter (ADC). The surface mount diodes employed are more compact than standard LEDs, and are more amenable to coating by solid-state sensor films. Results are presented from sensors employing a chemochromic ammonia sensitive film and demonstrate detection of this target in the ppm range. The configuration is applicable to a wide range of colorimetric based gas sensing materials.

IEEE Sensors Journal

This work may not be copied or reproduced in whole or in part for any commercial purpose. Permission to copy in whole or in part without payment of fee is granted for nonprofit educational and research purposes provided that all such whole or partial copies include the following: a notice that such copying is by permission of Mitsubishi Electric Research Laboratories, Inc.; an acknowledgment of the authors and individual contributions to the work; and all applicable portions of the copyright notice. Copying, reproduction, or republishing for any other purpose shall require a license with payment of fee to Mitsubishi Electric Research Laboratories, Inc. All rights reserved.

Copyright © Mitsubishi Electric Research Laboratories, Inc., 2005 201 Broadway, Cambridge, Massachusetts 02139



Low-Cost Surface Mount LED Gas Sensor

Roderick L. Shepherd, William S. Yerazunis, *Senior Member, IEEE*, King Tong Lau and Dermot Diamond*

Abstract— A low-cost chemical sensor comprising surface mount light emitting diodes (LEDs) has been developed for colorimetric based gas detection. The device consists of a pair of LEDs connected to a simple PIC microcontroller circuit, and in most basic form requires the use of only 2 I/O pins on the chip. The key features of this sensor are the use of an LED, rather than a photodiode for light detection, and an all-digital light detection protocol that leads to a reduction in cost and power consumption by avoiding the need for an Analog-to-Digital converter (ADC). The surface mount diodes employed are more compact than standard LEDs, and are more amenable to coating by solid-state sensor films. Results are presented from sensors employing a chemochromic ammonia sensitive film and demonstrate detection of this target in the ppm range. The configuration is applicable to a wide range of colorimetric based gas sensing materials.

Index Terms— LED sensor, gas sensor, ammonia sensor, light emitting diodes, colorimetry, optical sensor, low-cost sensor.

I. INTRODUCTION

EDs are being used far more commonly as light sources in optical chemical sensors due to the low-cost, low-power consumption, reliability and ever increasing range of devices and wavelengths available. The increased interest in LED sources has had a major impact on low-cost component based chemical sensors, where the main goal is to achieve analytical performance without the expense of more conventional instrumentation [1-5]. Typically a photodiode is used for detection, providing good sensitivity and a significant reduction in system cost. Usually the photodiode is operated at Vbias=0V and hence itself can be considered as a lowpower sensor, however, in addition to the detector, a good quality operational amplifier and mid-to-high resolution ADC are required to complete the device. These additional components not only increase system complexity and cost, but also add to the power requirements, which is of particular importance in battery-powered systems. The sensor we have developed requires only a simple microcontroller, and a pair of LEDs, one of which is driven in reverse bias mode to function

Manuscript received April 28, 2005. The financial support of the Adaptive Information Cluster (AIC) funded by Science Foundation Ireland (SFI03/IN3/1361) is gratefully acknowledged.

R. L. Shepherd, King Tong Lau, and D. Diamond are with the Adaptive Sensors Group, NCSR, School of Chemical Sciences, Dublin City University, Dublin 9, Ireland. (*corresponding author phone: +353-1-700-5404; fax: +353-1-700-8021; e-mail: dermot.diamond@dcu.ie).

W.S. Yerazunis is with MERL, Cambridge Research Lab, 201 Broadway, Cambridge MA 02139, USA.

as a light detector. The detector LED requires a short reverse biasing step (100µs) to the supply voltage (3V or 5V), but there are no other components requiring additional power so this is the only power draw associated with the detector itself. The system provides a simple, low-cost, battery powered sensing platform with very good sensitivity and signal to noise characteristics. Initially we developed this system for communication and light sensing applications that made use of the bi-direction properties of LEDs [6], after which we began work on all-LED colorimetric chemical sensors. To date we have reported on the use of a fused-LED sensor for monitoring colorimetric reactions in solution [7], and on the detection of Fe(II) at concentrations in the low ppb range using a pair of LEDs in transmittance mode [8]. Recently we presented results on a transmission mode solid state gas sensor for ammonia comprising two small surface mount LEDs and a chemochromic polymer sensing film [9]. This paper provides a more detailed account of the surface mount LED ammonia sensor, and outlines the key advantages of using this approach for colorimetric chemical sensing.

II. MATERIALS AND METHODS

A. Sensing Protocol

The sensing technique that we have developed is all-digital in nature, and hence there is no need for conversion of an analog signal as is normally the case with photodiode based optical chemical sensors. This reduces system cost by avoiding the need for a precision OPAMP and an external ADC chip or a more expensive microcontroller with a reasonably high-resolution inbuilt ADC. There is a short reverse biasing step (100us) associated with our LED sensor, which involves charging the diode capacitance to the supply voltage (5V or 3V), but this is the only power draw involved. While photodiode systems are typically operated at Vbias=0V, the power consumption of any supporting chips required for this method of detection (OPAMP and usually external ADC) will be significant. Hence the LED based sensing approach is low-cost with relatively low power requirements, and our findings thus far indicate that the signal integration approach used in the digitization of the photocurrent gives excellent s/n characteristics, and therefore very low limit of detection (LOD) in analytical measurements [8].

Typically, the detector LED is connected to a pair of digital I/O pins and operated in reverse bias mode. However, if pin availability it critical, the detector can be driven from a single

I/O pin with the anode tied to ground. The advantage of using a pair of I/O pins for detection is that the LED can be operated bi-directionally and hence can be used to emit and detect light. This has potential implications for sensors with communication capability, from simple visual feedback to more complex line-of-sight data transfer. In either case, the diode junction is charged to the supply voltage (normally 5 V), and then the I/O pin connected to the cathode is switched to high impedance input mode. Photons impinging on the detector diode produce a small (nA) photocurrent, which causes the 5V charge to decay. Using a timing routine implemented in firmware that is programmed onto the microcontroller, we continually poll the logic state of the input pin until the logic "0" threshold ($\sim 1.7 \text{ V}$) is reached. The time taken in microseconds to reach logic "0" is proportional to the amount of light detected. As the amount of light received increases, the diode discharges more rapidly, and hence lower timing values are observed. Conversely as the amount of light decreases the time to reach logic "0" increases. An oscilloscope trace of a sensing cycle (Fig. 1.) illustrates this process of diode discharge and logic threshold crossing. Logic voltage of the I/O pin is plotted on the vertical axis and time, in 200 microsecond divisions, along the horizontal axis. The time based measurement in this example is indicated by the horizontal line labeled 'count', and corresponds to a value of approximately 1300 µs. To increase noise immunity, the measurement process is not a simple timer that stops when the first logic "0" is read. Instead, repeated measurements are performed in rapid succession over the maximum time interval we wish to accept. For each measurement showing a logic "1", we increment a counter; if the measurement shows a logic "0" we do not change the counter. Assuming that the noise signal is uncorrelated with the timing of the sample measurements, this gives us a significant improvement in S/N ratio, and also yields a constant and deterministic output rate for the sensor/microcontroller system.



Fig. 1. Oscilloscope trace of LED sensor discharge curve.

Digital detection of light is achieved by continually polling the I/O pin logic state. At present data is collected via an RS232 link between the device and a PC, but this could easily be replaced with an LED based wireless communication link.

The circuit employed for this work (Fig. 2) comprised a PIC16F876 microcontroller, MAX232ACP RS232 level shifter and supporting components required by the chips, including a 20MHz ceramic resonator for the PIC. The microcontroller has an inbuilt 10-bit ADC, but this feature is not required for our sensor, and hence was unused. While many reasonably priced microcontrollers such as the PIC16F876 now have an internal ADC, currently these tend to be more general-purpose converters where the resolution is only 10-bit. In many sensing applications better resolution is required, so typically a 12-bit or even 16-bit external ADC is employed, adding significant cost to the system and increasing power usage. The current PIC16F876 based circuitry provides a generic interface and control platform that is used for the development of a range of different sensors in our labs, but the sensing technique described herein could easily be implemented on a lower-cost microcontroller without an ADC. The 16F876 is running at 5 MIPS, and the resulting minimum sensing resolution achievable at this clock speed is approximately 5 µs. The maximum fixed sensing period typically employed is $65000 \ \mu s$, which ensures that a suitable integration time is provided without causing any overflow problems (timing values are stored in a 16-bit variable, max. = 65535). Significantly longer integration periods are possible by adding an additional delay in the timing routine implemented in software on the PIC. This scaling permits measurements to be made over hundreds of milliseconds or even seconds if required, but there is an obvious trade-off between the total integration time and resolution, which decreases with increased scaling.



Fig. 2. Block diagram of PIC16F876 LED sensing circuit.

If noise is considered (typically $\pm 10 \ \mu s$ on the observed discharge time) then the minimum sensing resolution is closer to 40 μs , but this still provides more than adequate sensitivity for the measurements described in this work.

B. Sensor Design

The sensor comprises two 1206 format surface mount LEDs, which have overall dimensions of 3.2 x 1.6 mm and a flat lens (Fig. 3). For the sensor described in this paper, a 574 nm green exciter LED (Kingbright KP-3216MGC) was used in combination with a 621 nm red-orange detector LED (Lite-On LTST-C150KAKT). The LEDs are soldered to Molex 2.0 mm pitch header pins for ease of connection and to allow for adjustment of the air-gap (Fig. 3b, Fig. 4). A variety of arrangements are possible, but for this paper we focused on the design shown in Fig. 4, where the diodes are stacked vertically with respect to each other. For this work the detector LED, which is coated with the chemochromic reagent film, was positioned such that it faced downwards, effectively reducing effects arising from variations in ambient light. In particular, this arrangement ensures that the majority of room lighting does not impinge directly on the detector diode. The exciter LED, which was used without any additional coating on the lens, was positioned directly below the detector at a distance of approximately 0.5 - 1.0 mm. The LEDs were plugged into a 2.0 mm Molex receptacle, and the air-gap between the LEDs adjusted by simply moving the detector LED relative to the exciter (Fig. 4). This physical adjustment of the LEDs with respect to each other allows for coarse tuning of the baseline signal, which is particularly useful when relocation of a device involves major changes in ambient light levels.



Fig. 3. Schematic of the surface mount LED ammonia sensor illustrating the form factor of the 1206 surface mount LEDs (a) and configuration of LEDs in the chemical sensor (b).

C. Ammonia SpecificCcolorimetric Reagent

The reagent used for this work is p-Nitrophenylnitrosamine (NPNA), and was synthesized in-house (Fig. 5). A mix of NPNA and PVC in cyclohexanone was prepared, which was subsequently drop coated onto the detector LED lens by applying a small volume from a pipette. Due to the relatively low surface area of the LEDs used in this work, the volume of reagent required for preparation of each sensor was kept to a minimum (< 5 μ L). NPNA forms a weak complex with ammonia (Fig. 5) and causes a chromic shift in the material from yellow to red. NPNA is pH sensitive to some extent, but only extremely basic species shift the equilibrium. As there are limited cross-reacting species in the gas phase with these properties, NPNA in the solid state exhibits reasonably high selectivity to NH₃ and primary amines.



Fig. 4. Surface mount LED chemical sensor housed in molex receptacle. Coarse baseline adjustment is made by changing the air-gap between LEDs.

D. Ammonia SensorTesting

The gas sensor was tested for responses to NH_3 in a simple flow cell that was fabricated in-house. A 100 ppm (v/v) NH_3 standard premixed gas cylinder (Scott UK) was used in combination with a nitrogen line for dilution to deliver ammonia samples in the concentration range 6.25 - 14.3 ppm to the sensor. Gas flow rates were controlled by rotameters, and were in the range 0 - 50 mL/min for NH_3 and a constant 300 mL/min for N2. The nitrogen flow was maintained at 300 mL/min throughout all experiments, both during response and recovery cycles, as the manual adjustment on the rotameters used was not precise enough to allow for continual flow adjustments. The gas flow was passed through the headspace of a water bubbler to maintain relatively constant humidity throughout the experiments.

III. RESULTS AND DISCUSSION

A. NPNA Optical Response

The optical response of NPNA upon exposure to ammonia is due to shifts of the protonated form absorbance peak (λ max = 420 nm) to the deprotonated basic form ($\lambda max = 550 \text{ nm}$) (Fig. 6). This effects a visible colour change in the material, which is yellow in air and red after interaction with ammonia. The green emitter LED chosen for this particular sensor has a peak wavelength of 574 nm, a dominant wavelength of 568 nm and a spectral half-width of approximately 20 nm [9]. The emission spectrum of the LED was measured to confirm these specifications using an Ocean Optics S2000 spectrometer, and for comparative purposes has been superimposed over the NPNA spectra in Fig. 6. It is clear that the emissive properties of this LED overlap well with the base form of NPNA, and hence make it an ideal light source for monitoring the development of the red NPNA/ NH₃ complex. As the concentration of NH₃ increases, the intensity of light transmitted through the sensing film to the detector LED decreases, and the discharge time values measured by the microcontroller circuit increase.



Fig. 5. Reaction scheme of NPNA with NH₃. Ammonia forms a weak complex with NPNA and this brings about a colour change from yellow to red.

B. LED Sensor Responses

The sensor was exposed to NH₃ samples ranging from 6.25 to 14.3 ppm. As was mentioned earlier, an approximate lower sensing limit of this device based upon observations of the baseline noise is around 20 µs of dither in the measured nominal discharge time, so at the lower concentrations of NH₃ tested the sensor was approaching the limit of detection. Sensor responses to multiple samples of 9 ppm NH₃ are shown in Fig. 7, and as anticipated, increases in the sensor timing response were observed in the presence of ammonia. The level of noise present in the signal is quite significant, but despite this the responses are clearly distinguishable and reproducible. By applying a simple moving average filter to data (Microsoft Excel, period = 20) there is a significant improvement in the s/n ratio (Fig. 7 solid line). Some baseline drift was observed (Fig. 7), which can be attributed to equilibration of RH in the sensor test chamber over the course of the experiment. The drift is most evident at the start of the



Fig. 6. Spectra of NPNA acid and base forms, and emission spectrum of green 574nm emitter LED. The overlap between the LED source and base form of NPNA, which develops in the presence of ammonia, is very good.

test-run (peak 1) with the baseline virtually stabilising by the last sample (peak 4). Experiments were carried out at essentially saturated RH, which provided a simple means of maintaining humidity close to constant, but caused some drift due to polymer swelling as the RH stabilised. Despite some baseline instability sensor responses were not significantly affected and quite reasonable reproducibility was observed from one sample to the next.



Fig. 7. Response of LED sensor to multiple samples of 9 ppm NH3 in nitrogen. Raw data is plotted in grey, and the smoothed response in black after a moving average filter (per = 20) was applied to the raw data. Increases in the sensor timing response are observed in the presence of ammonia due to stronger absorbance of emitted light by the NPNA film.

The simple flow system used to prepare and deliver these gas samples did not allow precise control of the NH₃ flow rates, and the influence of this on sensor reproducibility became more evident at faster rates where the ammonia accounted for a larger fraction of the total flow. The larger errors associated with the 12.5 and 14.3 ppm data points (Fig. 8) are indicative of this flow induced variation in response. Despite these sources of variation, responses were linear over the entire concentration range investigated as shown in Fig. 8. The NH₃ range studied was quite limited, as we were primarily concerned with determining or atleast estimating the limit of detection (LOD). The minimum NH₃ concentration that could be generated with the set-up used was 6.25 ppm, and hence it was not possible to obtain data at lower concentrations.

The estimated LOD, which is based on baseline noise of the raw data and an average response at 6.25 ppm NH₃ of approximately 30 us, is about 5 ppm in this particular configuration and experimental design. We are confident that this can be significantly improved, by applying simple filtering as discussed above, and by varying the polymer sensing layer thickness, which has a significant effect on both response time and sensitivity. Until now we have coated the detector LED in our sensors by simply drop coating a small volume of NPNA/polymer onto the lens and allowing this to dry. While this has been adequate for initial prototype sensors, the ideal approach to controlling the film thickness is spin coating. The surface mount LEDs used are ideal for use with this deposition technique, as the lens structure provides a uniform flat surface. Controlling the volume of sensing material and rotation rate will provide a means of varying the NPNA film thickness, enabling further optimisation of the sensors. We plan to investigate this further in the future, but are encouraged by the results already obtained using the sensor described in this work. The current surface mount LED sensor, which is clearly not optimised, detected ammonia in a linear fashion in the 5 -15 ppm range.



Fig. 8. Average changes in response (n = 4) of LED ammonia sensor with error bars indicating the standard deviation.

C. System Flexibility and Portability

One of the key features of our system, is the software based timing protocol, and in addition to the aforementioned benefits of the digital signal this provides, many sensing parameters can be controlled and updated in software. Critical sensing properties including integration period, minimum timing resolution, average rate, sampling rate, and power saving schemes can easily be controlled at run-time in code or updated by an external device without any change to the physical characteristics of the sensor itself. If the detector LED is configured to operate bi-directionally, these sensor properties can even be updated using LED-to-LED wireless communication.

The sensors are also highly portable, again due to the software based technique used to measure detector photocurrent, and are therefore effectively platform independent. LED based sensors can be interfaced seamlessly with a wide range of different microcontroller architectures and microcontroller-based devices. The main task when moving between different processors involves porting a small piece of firmware, and this could be streamlined by developing a library of drivers to support the more common parts. Potentially LED based sensors can be incorporated into a diverse range of devices, but one area of major significance is the emerging field of wireless sensor networks (WSN). These systems are battery powered, microcontroller based communication and sensing nodes that rely on low-power components and highly efficient power management in order to extend lifetime. Various platforms are being developed that are based on a range of different microcontrollers [11 - 14]. In all cases there are versions of these platforms that run off 3V lithium coin cells, and these batteries have small energy capacities (typically 200 - 600 mAh at 3 volts). Power efficiency is therefore paramount, and saving a few mA of current can significantly increase the lifetime of an individual node in the network. These LED-based sensors are ideally suited for hardware platforms such as these with limited power, and we are currently investigating the integration of our sensors with low-power wireless nodes. The ultimate goal is to deploy low-power WSNs for environmental and industrial monitoring, with the sensing tasks performed by low-cost LED chemical sensors.

IV. CONCLUSIONS

A low-cost colorimetric ammonia gas sensor has been developed that uses small surface mount light emitting diodes as both light source and detector. Sensor responses are monitored using a simple microcontroller circuit, which is entirely digital in nature. This configuration avoids the cost and power requirements of typical supporting components employed when using a photodiode (precision OPAMP and expensive external ADC/more expensive microcontroller with high-resolution internal ADC), and as the sensing protocol employs a software based photocurrent measurement, the devices are highly flexible and essentially platform independent. The sensor has been demonstrated for the detection of low ppm levels of ammonia, as determined using a selective colorimetric polymer coating containing NPNA, but this configuration is amenable to a wide range of colorimetric based gas sensing materials. Indeed virtually any colorimetric assay could be immobilized in some way and deposited onto the lens of a surface mount LED. The challenge is really in developing reliable sensing materials for this purpose, with the LED sensor providing a low-cost generic platform upon which sensors can be developed. The portability, low-cost and lowpower characteristics of these devices make them ideal chemical sensors for WSN applications where power consumption is critical.

REFERENCES

- K.T. Lau, S. Edwards and D. Diamond, "Solid-state ammonia sensor based on Berthelot's reaction" Sensors and Actuators B, vol. 98, pp. 12-17, 2004.
- [2] L. Byrne, K.T. Lau, and D. Diamond, "Monitoring of headspace total volatile basic nitrogen from selected fish species using reflectance spectroscopic measurements of pH sensitive films", The Analyst, vol. 127, pp. 1338-1341, 2002.
- [3] Y.-S. Lee, B.-S. Joo, N.-J. Choi, J.-O. Lim, J.-S. Huh, and D.-D. Lee, "Visible optical sensing of ammonia based on polyaniline film", Sensors and Actuators B, vol. 93, pp. 148-152, 2003.
- [4] E.J. Cho and F.V. Bright, "Intergrated chemical sensor array platform based on light emitting diode, xerogel-derived sensor elements, and high-speed pin printing", Analytica Chimica Acta, vol. 470, pp. 101-110, 2002.
- [5] F.A.A Matias, M.M.D.C. Vila and M. Tubino, "A simple device for quantitative colorimetric diffuse reflectance measurements", Sensors and Actuators B, vol 88, pp. 60 – 66, 2003.
- [6] P. Dietz, W. Yerazunis, D. Leigh, "Very Low-Cost Sensing and Communication Using Bidirectional LEDs", UbiComp 2003: Proceedings, vol. 2864, pp. 175-191, 2003.
- [7] K. Lau, S. Baldwin, R. Shepherd, P. Dietz, W. Yerazunis and D. Diamond, "Novel fused-LEDs devices as optical sensors for colorimetric analysis", Talanta, vol. 63, pp. 167-173, 2004.
- [8] S. Baldwin, K.T. Lau, R. L. Shepherd, W.S. Yerazunis and D. Diamond, "Colorimetric Detection of Iron(II) Using Novel Paired Emitter Detector Diode (PEDD) Based Optical System", IEICE TRANS. ELECTRON., Vol. E87-C (12) pp. 2099 – 2102, 2004.
- [9] R. L. Shepherd, W. S. Yerazunis, K. T. Lau, and D. Diamond, "Novel Surface Mount LED Ammonia Sensors", Proceedings IEEE Sensors 2004, pp. 951 – 954, 2004.
- [10] Kingbright KP-3216MGC Technical Specification SPEC NO:DSAB1394, www.kingbright.com, 2002.
- [11] http://www.xbow.com/Products/Product_pdf_files/Wireless_pdf/MICA 2DOT Datasheet.pdf
- [12] http://www.tyndall.ie/research/mai-group/25cube mai.html
- [13] http://www.smart-its.org/artefacts/artefacts.html
- [14] http://www.doc.ic.ac.uk/vip/ubimon/bsn node/index.html

Roderick L. Shepherd received his BEnvSc (Hons) and PhD degrees in 1996 and 2001, respectively, from the University of Wollongong, Australia. He worked as a research fellow in the National Centre for Sensor Research (NCSR), Dublin City University 2001 – 2002, and then as a visiting research scientist in Mitsubishi Electric Research Laboratories (MERL), Cambridge Massachusetts, USA in 2003. He is currently a research fellow in the Adaptive Sensors Group, NCSR at Dublin City University working primarily on wireless sensor networks and low-cost LED based optical chemical sensors. His research interests include chemical sensors, inherently conducting polymers, colorimetric optical sensors, robotics and wireless communication.

William Yerazunis has worked in a number of fields including: optics, vision processing, and signal processing (for General Electric's jet engine manufacturing); computer graphics (at Rensselaer's Centre for Interactive Computer Graphics); artificial intelligence and parallel symbolic computation (for DEC's OPS5, XCON, and the successor products such as RuleWorks); radioastronomy and SETI (at Harvard University), transplant immunology (for the American Red Cross), virtual and augmented reality (Diamond Park and SPLINE), real-time sensing and ubiquitous computing (iGlassware), real-time statistical categorization of texts (the CRM114 Discriminator anti-spam system) and LED-based chemo-physical sensors and communication systems. He has appeared on numerous educational television shows, and holds 22 U.S. patents.

King Tong Lau is currently a senior research chemist in the National Centre for Sensor Research, Dublin City University. He received his PhD from University of London (2001) on electrochemical biosensor development. His research interest is in the area of molecular recognition and development of low cost, deployable sensor devices for chemical, environmental and food quality monitoring.

Dermot Diamond received his Ph.D. from Queen's University Belfast (Chemical Sensors, 1987), and is currently the PI of a recently awarded Science Foundation Ireland Grant (SFI) and director of the Adaptive Sensors Group, NCSR, Dublin City University. The Adaptive Information Cluster is an SFI funded initiative that brings together DCU and UCD. The AIC's goal is "to capture the world digitally in a comprehendible way". As such, the AIC is integrating leading-edge research in wireless sensing, content extraction from video and audio media, collaborative reasoning and personalization. He has published over 100 peer-reviewed papers in international science journals, and is co-author and editor of two books both published by Wiley. His research interests range from molecular recognition, host-guest chemistry, ligand design and synthesis, electrochemical and optical chemical sensors and biosensors, lab-on-a-chip, sensor applications in environmental, clinical, food quality and process monitoring, development of fully autonomous sensing devices, wireless sensors and sensor networking.