

# Learning to select distinctive landmarks for mobile robot navigation

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## Abstract

In landmark-based navigation systems for mobile robots, sensory perceptions (e.g., laser or sonar scans) are used to identify the robot's current location or to construct internal representations, maps, of the robot's environment. Being based on an external frame of reference (which is not subject to incorrigible drift errors such as those occurring in odometry-based systems), landmark-based robot navigation systems are now widely used in mobile robot applications.

The problem that has attracted most attention to date in landmark-based navigation research is the question of how to deal with perceptual aliasing, i.e., perceptual ambiguities. In contrast, what constitutes a good landmark, or how to select landmarks for mapping, is still an open research topic. The usual method of landmark selection is to map perceptions at *regular* intervals, which has the drawback of being inefficient and possibly missing 'good' landmarks that lie between sampling points.

In this paper, we present an automatic landmark selection algorithm that allows a mobile robot to select conspicuous landmarks from a continuous stream of sensory perceptions, without any pre-installed knowledge or human intervention during the selection process. This algorithm can be used to make mapping mechanisms more efficient and reliable. Experimental results obtained with two different mobile robots in a range of environments are presented and analysed. © 2001 Published by Elsevier Science B.V.

*Keywords:* Landmark-based navigation; Mobile robots; Kalman filter; Automated landmark selection

## 1. Introduction

### 1.1. General considerations — approach

For mobile robot navigation over realistic distances, navigation systems based on sensory perceptions or landmarks are usually used. This is because, unlike dead-reckoning methods, perception-based systems

do not suffer from drift error. However, they are prone to the problem of perceptual aliasing, where a number of perceptions from different parts of an environment look similar.

There is another very relevant aspect of landmark-based navigation that has not yet been solved satisfactorily. This is the question of what constitutes a 'good' landmark, i.e., the question of how landmarks should be selected. So far, this problem has largely been ignored by either logging sensory perceptions at regular intervals, or comparing them with pre-installed models of 'good' landmarks supplied

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by a human supervisor. The first method is unsatisfactory because of its inefficiency, while the latter has the risk that a human supervisor will either select landmarks that are not easily recognisable by the robot, or overlook landmarks that are inconspicuous to a human, yet are perfectly recognisable by a robot.

In this paper, we present an algorithm that addresses this question of landmark selection, and facilitates the automatic selection of landmarks, i.e., without any human supervision. The algorithm allows a mobile robot to select landmarks for navigation from a continuous stream of perceptions, without using any external (user-supplied) feedback. The principal assumption is that a landmark that is suitable for navigation should be both conspicuous to the robot's sensors and reliably detectable. This paper first describes the landmark selection algorithm, which selects conspicuous landmarks. It then introduces performance measures that evaluate the consistency of landmark selection, and applies them to the experimental results obtained with two different mobile robots in a range of real-world environments.

### 1.2. Method of landmark selection

In this paper, landmarks are selected by defining conspicuous landmarks to be those that are unexpected, i.e., perceptions that do not conform to an acquired model of the temporal relationship between successive sensory perceptions. The landmark selection system described in this paper uses a neural network to learn a general model of the relationship between a robot's successive sensory perceptions, acquired over a range of different environments during a training phase. This general model is then used to predict future sensor readings based on current sensor readings. Any perception where the prediction does not match the actual reading obtained at the next time step to some accuracy is selected as a landmark on the grounds that it differs markedly from the usual relationship between successive perceptions — it is conspicuous.

Two methods of selecting those places where the prediction fails to predict the next perception accurately are proposed, one based on detecting peaks in the prediction error curve, and the other using a Kalman filter to maintain a model of the error and

selecting as landmarks those places that fall beyond the error bounds of the Kalman filter.

### 1.3. Performance evaluation

A very important feature of a landmark is that it can be *consistently* detected on every run, so that the robot can navigate using the landmarks. For this reason, a metric is proposed by which the consistency of the landmark selection can be evaluated and the various detection techniques compared. This is discussed in Section 5.

### 1.4. Key contributions

The experiments discussed in this paper demonstrate that it is possible to devise algorithms that enable a mobile robot to select consistent landmarks for navigation automatically. User guidance or pre-installation of models is not required. One fundamental feature of the approach presented in this paper is that a model of the temporal relationship between sensory perceptions is used, i.e., the robot is *robot-centred*, rather than *environment-centred*. This means that the algorithm is able to select landmarks in any environment — even completely novel ones — after training.

The experiments have been conducted on two different mobile robots, in a range of different environments. The results demonstrate that the algorithm is not sensitive to hardware aspects, nor properties of a specific environment. In order to assess the consistency of landmark selection, two measures of performance are introduced: the sum-squared-difference between prediction errors of different traversals of the same environment, and the landmark consistency measure LMC, which evaluates whether the algorithm selects the same location as a landmark during each traversal of an environment.

## 2. Related work

A number of methods of selecting landmarks are described in the literature. Some systems [3,15] take sonar scans at regular intervals, such as every 150 cm of travel in an environment, with every sonar scan being used as a landmark that is put into the map. This avoids the problem of ensuring that landmarks should

be detected consistently, but since in many environments — such as corridors — a large number of perceptions are almost identical, the map is filled up with information that does not aid navigation.

Another problem with mapping at regular intervals is that perceptions that lie between sampling locations are missed. It may well be that these missed perceptions include considerably more distinctive landmarks than those perceived at the user-defined sampling locations.

A number of researchers have considered selecting suitable landmarks for mobile robot navigation. These can be split into off-line approaches that are not suitable for use on the robot as it explores, and systems that can be used on-line.

In the first category is the technique of asking the user to define the landmarks before the robot explores. Humans typically select objects that they find easy to recognise, such as doors, or line segments extracted from camera images recorded as the robot travels [6]. These approaches suffer from the same problem — the robot's perceptions of the world are very different to those of a human, and so features that the researcher thinks are distinctive may not be noticeable by the robot.

In contrast to this, Thrun [13] addressed the problem of Bayesian learning to select an optimal set of landmarks for performing self-localisation in one specific environment, where the landmarks consisted of a projection from the robot's raw sensory perceptions (camera images) onto vectors in a lower-dimensional space. This optimisation was performed by minimising directly the quantity of interest, namely the robot's error in self-localisation. Thrun showed that his method produced better performance than localisation using designer-determined landmarks including doors and ceiling lights. A similar, but computationally cheaper technique, was developed by Vlassis et al. [14], who showed that their optimisation method produced better results than principal component analysis. Our approach differs in that we do not carry out any analysis of the utility of the landmarks selected, but instead use a self-acquired model of 'typical' sequences of perceptions, which is independent of any particular task or environment.

Neither of the above approaches are suitable for *on-line* selection of landmarks, because they rely on an acquired model of a previously explored environ-

ment. Zimmer [16] considered the problem of selecting landmarks in a topological map through a process of 'life-long learning', where the robot's map was continuously adapted on-line during exploration. This approach used global statistical information, based on comparison of accumulated error statistics at each of the nodes, to decide where to add and delete nodes in the map.

A related idea can be found in Bourque and Dudek [2], who addressed the 'vacation snapshot' problem of deciding in which locations to take camera images in order to obtain a set of images that best represent an entire environment. This approach kept running statistics on what is a 'typical' perception, together with backtracking to previously visited locations that were subsequently found to be 'atypical'. Our method differs from these approaches in that we only use *local* sensory information to decide when to add landmarks to the map.

Simhon and Dudek [12] also use only local information. They addressed the problem of deciding the best locations in which to create local metric maps, using a distinctiveness function based on the robot's ability to localise itself within such a map. However, this approach relies heavily on the designer's intuition in selecting an appropriate distinctiveness function.

### 3. Acquiring a general model of the relationship between consecutive sensory perceptions

The first part of the proposed system for selecting landmarks is the general model of the relationship between consecutive sensory perceptions of the robot. The perceptions are sampled at regular spatial intervals as the robot travels, using dead reckoning to estimate the distance between samples.

One way for the robot to acquire the model is through the use of a neural network trained on data collected by the robot during exploration. Since the model should predict the next perception that the robot will encounter, a supervised learning technique is suitable, where each input vector is paired with a target output (in this case the sensory perception perceived at the next time step), and the weights of the network adjusted to reduce the error in the prediction, usually by a gradient descent learning method.

### 3.1. The neural network architecture

The most simple supervised network is the perceptron, a single-layer feedforward network [1,11]. The inputs to the network are directly connected to the outputs by adjustable weights, and the update rule modifies these weights. More complex networks have intervening layers of ‘hidden’ units, and the difference between the network output and the target is propagated back through the network to adapt the strength of the connections between the units.

In the current application we do not desire a complex model of the perceptions. Rather, a very simple model that contains only the most general features is useful, so that there are environmental features for which the prediction is not good, since poor prediction is what generates the landmarks. For this reason, a perceptron is used rather than any more complicated model. This network calculates the output  $o_k$  for each output unit  $k$  at the current time using Eq. (1):

$$o_k = f(\mathbf{w}_k \cdot \mathbf{i}), \quad (1)$$

where  $\mathbf{i}$  is the input vector,  $\mathbf{w}_k$  is the weight vector of output unit  $k$ , and  $f$  the transfer function. In our experiments this was the sigmoid function given in Eq. (2),

$$f(x) = \frac{1}{1 + e^{-2x}}. \quad (2)$$

The network weights are updated during training using Eq. (3):

$$\mathbf{w}_k(t+1) = \mathbf{w}_k(t) + \eta(\tau_k - o_k)\mathbf{i}, \quad (3)$$

where  $\tau_k$  is the target value for output unit  $k$ , and  $\eta$  is the learning rate (0.2 in the experiments reported here).

#### 3.1.1. Modifying the connectivity

As the prediction that is being made is of future sensory perceptions, and the robot is always driving forwards, the future perceptions of any one sensor depend only on the sensors in front of it. The network connectivity was chosen to model this fact, so the perceptions of sensors facing forwards are used as inputs to model rear-mounted sensors, but not vice versa. The reading of a sensor at the current time is also used in the prediction of its own next reading. The architecture is shown in Fig. 1 and is referred to in this paper as the ‘modified perceptron’. In addition to the network inputs shown in the figure, a bias input that is permanently set to  $-1$  is used.

#### 3.1.2. Adding temporal inputs

The modified perceptron introduced in the previous sections learns to predict the next reading given the current one. Another possibility for modelling the temporal relationship between sensory perceptions

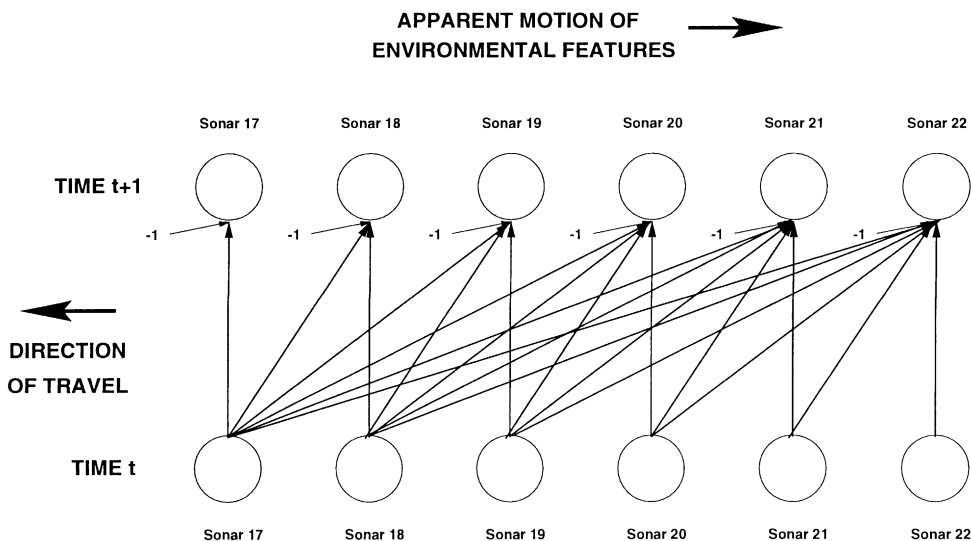


Fig. 1. The neural network architecture of the modified perceptron. All connections are shown. A bias input, permanently set to  $-1$ , which was connected to all output units was also used.

would be to use information from past perceptions. Two methods of doing this are investigated in this paper.

The first is the lagged perceptron, shown in Fig. 2. This network uses a lagged input vector, so that the inputs to the perceptron at the current time consist not just of the current readings, but also the readings of  $n$  previous time steps.

The second technique, shown in Fig. 3, is the recurrent perceptron. In this network, the inputs to the network are the outputs at previous time steps (as well as the current perception), so there is a recurrent mapping between the outputs at the current time and the inputs at the next time.

In both cases, the weight update rule described in Eq. (3) was used in exactly the same way as before, and the bias input was still present. For both of these networks, a value of  $n = 1$  was used, so that only one previous time step was considered. Preliminary investigations showed that this generalised the best between different environments.

### 3.2. Sensor signal pre-processing

The sonar readings that are used as inputs to the neural network are intrinsically very noisy. Typical problems include cross-talk, where a sonar pulse emitted by one sensor is received by another and mistaken as its own, and specular reflection, where a sonar pulse rebounds from a surface away from the robot and therefore suggests that objects are much further away than they actually are. This results in a spike in the sonar readings.

One way to deal with this problem would be to ignore any sonar response that differs from the previous one by too much. However, this would be a problem when the robot went past an open doorway, when there would, correctly, be a large discontinuity in the readings. Instead, a median filter can be used. This filter replaces the value at each time  $t$ , with the median average of the previous  $k$  time steps. Since the door is wide enough to appear on several readings, it will pass the filter.

In the experiments reported here,  $k$  was chosen to be 5, so that for each sensor input, the median of the readings for that sensor at times  $t, t - 1, \dots, t - 4$  replaced the reading at time  $t$ . The problem with using the median filter is that it blurs the edge be-

tween the readings before and after the discontinuity, delaying the perception of the feature. However, in the present application this is not a problem, as the same effect will occur each time the robot passes that place and so the perception will be the same on each occasion.

### 3.3. Network training procedure

One factor that affects neural network performance is the amount of training that is done. For the type of behaviour that the model should exhibit, i.e., a generalised prediction that is not specific to any individual environment, this is especially important. In this work, the technique of early stopping [1] was used to decide how many learning iterations should be applied. Early stopping monitors the error of the network, evaluated on an independent validation set, stopping the training of the network when the prediction error on the validation set stops decreasing. This suggested that about 20 epochs, or complete runs through the data, were sufficient with the learning rate of  $\eta = 0.2$  that were used. The data that were used for this purpose were collected from 10 runs through an environment. Five of the runs were used as training data and the other five as validation data. Once the number of epochs required were found, the data were discarded.

Another important problem is to decide how big the training set should be, i.e., how many environments should make up the training set, and how many runs from each environment should be used. The second question was not considered to be a problem. A total of 20 runs were collected in each environment, and these were split into two sets of 10, with one set for training and one set for testing. Since the network should learn a general model of the entire environment in which the robot will operate, it makes sense that the network should be trained by using examples from a large number of the different environments that the robot can possibly encounter in operation. For this reason, in the results presented here two different types of training set were considered. In the first, data from three different environments in which data were collected were selected, with different combinations of environments used, while in the second, data from all of the environments were used.

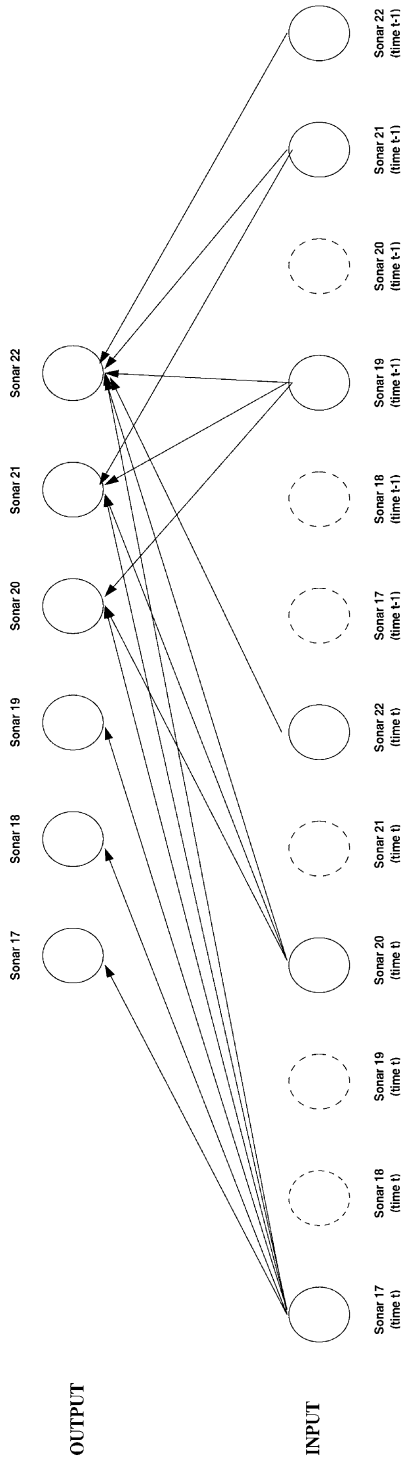


Fig. 2. The lagged perceptron. The inputs to the network comprise the sonar readings at times  $t$  and  $t - 1$ , with the reduced connectivity used for both sets of inputs. For clarity, the connections are only shown for a subset of the neurons.

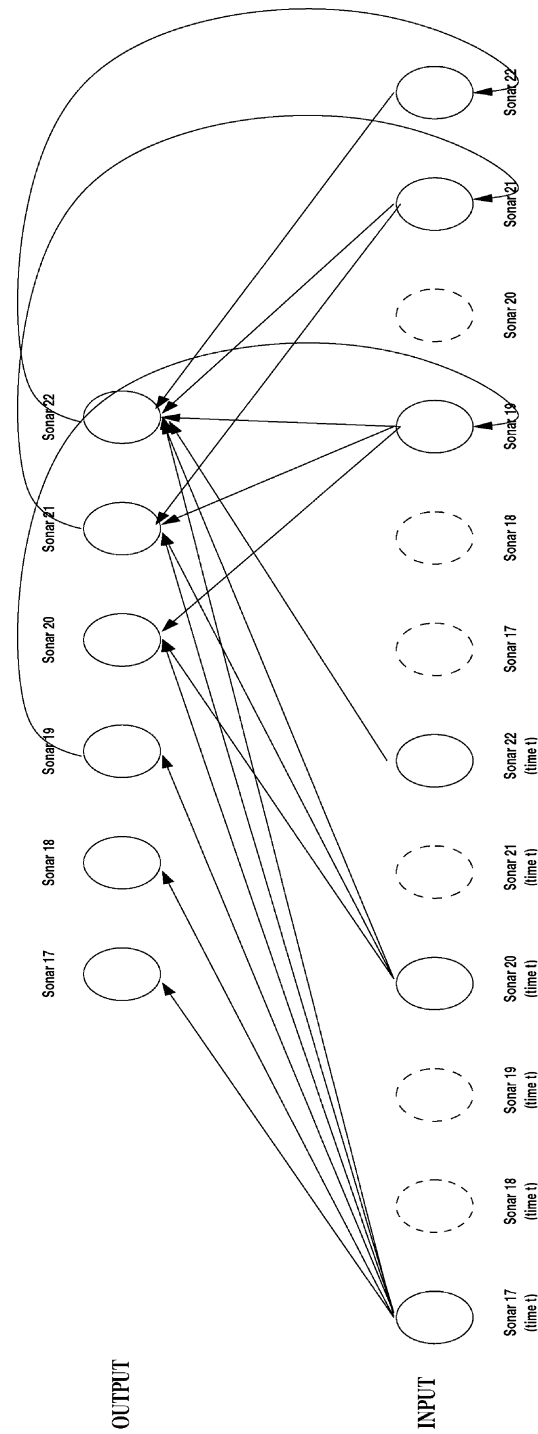


Fig. 3. The recurrent perceptron. The outputs of the network, the predicted perception at the previous time step, are used as inputs as well as the current perceptions of the robot. Again, for clarity, the connections for only some of the neurons are shown.

#### 4. Selecting landmarks with the trained network

Once the neural network has been trained, the current perceptions can be fed into the model to produce a prediction of the next perception. This is done by calculating the output value,  $o$ , using Eq. (1). This output is then used to compute the error,  $F_1$ , at the current time, by computing the squared difference between the predicted output and the actual perception obtained at time  $t + 1$ .

In the landmark selection technique presented here, the places where the prediction of the general model is bad — the places where the error is high — are used as landmarks. The ‘quality criterion’ for suitable landmarks here is that they are selected *consistently*, and the hypothesis is that the places where the model breaks down (resulting in bad predictions) will be the same on every run. The problem is to detect these peaks in the error curve that are above the level of noise in the data. Two ways of doing this are described below.

##### 4.1. Selection method 1: peak detection

The first method of detecting locations where prediction is bad is to detect peaks in the error curve. There are a number of ways in which peaks can be found. The problem is that the error curve is very noisy, as can be seen in the typical error curve shown in Fig. 4. Using a perceptron means that the accuracy of the predictions is never particularly high, so only peaks that stand out above the residual noise level should be detected.

The peak detection itself can be done by looking for zero crossings of the first derivative. This is approximated by considering points where the sign of  $\Delta t / \Delta F_1$  changes, where  $F_1$  is defined as

$$F_1(t) = \sum_j [o_j(t-1) - i_j(t)]^2 \quad (4)$$

with  $o_j(t-1)$  being the output of network unit  $j$  at the previous time step and  $i_j(t)$  the perception of the equivalent robot sensor at the current time step.

A typical error curve is shown in Fig. 4. It can be seen that, while there are a number of significant peaks in the error curve, in addition there are several very minor peaks within the noise level of the data.

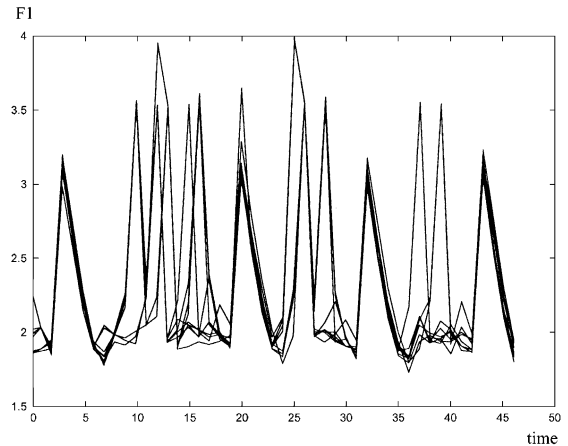


Fig. 4. Typical output error curves over several runs in the same environment. The way the error occasionally rises above a baseline can be seen, as can the way the different curves match well at some points, but less so at others.

These will also be detected by using the peak detection algorithm, which is undesirable. For this reason, a number of filtering steps were carried out before the peak detection. These filters are designed to smooth the curve, reducing the number of peaks, so that only those that are significant stand out. The process that is performed is shown in Fig. 5 and is described below. More detailed analysis of the effects of the results are given in [4,9].

- *Integration.* The current error value,  $F_1(t)$ , was integrated with the previous nine, effectively providing a mean average of the 10 values.

$$F_2(t) = \sum_{i=0}^9 F_1(t-i). \quad (5)$$

- *Median filter.* The last nine integrated values were median filtered in order to remove surface noise from the curve. This provides a value  $F_3(t)$  for each time  $t$ .
- *Smoothing.* The curve was then smoothed, removing any remaining small peaks by using a momentum term from previous time steps (Eq. (6)).

$$F_4(t) = \lambda F_3(t) + (1-\lambda)F_4(t-1), \quad (6)$$

where a value of  $\lambda = 0.2$  was used. This means that the filter uses information about the previous value to smooth out the curve. The drawback of this

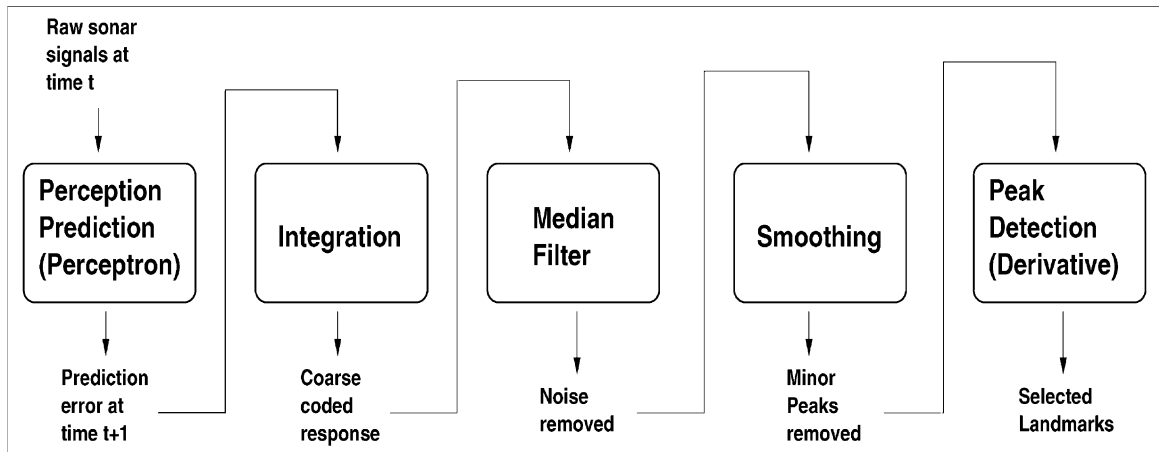


Fig. 5. The filtering mechanism applied before peak detection.

technique is that it also pushes the position of the peak back, so that the robot does not perceive the peak in the place where the readings were taken, but some way further along.

#### 4.2. Selection method 2: Kalman filtering

The filters described in Section 4.1 have the side-effect of delaying the appearance of peaks in the data and smearing them across the data. For this reason, an alternative approach was considered, using a Kalman filter [5] to maintain a model of the error curve and look for places where the error did not fit the model.

The Kalman filter was designed to produce successively better approximations to the state of a linear process that is controlled by a difference equation. It works by maintaining a continuously updated estimate of the state of the process being measured (here, the predicted sensory perceptions), including an estimate of the error in the measurements. By using the filter to keep a model of the error curve and detecting when the predictions of the model are beyond error bounds, outliers to the data at the current time can be detected. These outliers are suitable candidates for landmarks. For further details, see [7].

The implementation used in this work is a very simple Kalman filter. A more complete description is given in [8]. The Kalman filter maintains an estimate of the covariance of the data (in this simple,

one-dimensional case, the variance,  $v$ ) and uses that to put error bounds on the predictions, using the Kalman gain,  $K$ . At each time step, a number of equations are applied. The first, Eq. (7), calculates the Kalman gain for the new iteration, based on the estimate of the variance at the previous time step.  $R$  is an estimate of the variance in the measurement error, i.e., the error that is introduced in the reading of the state. Then, in Eq. (8), the new estimate of the state of the predicted sensory perception,  $y(t)$ , is calculated. This includes a (noisy) measurement of the predicted sensory perception taken at the current time,  $x(t)$ . Finally, Eq. (9) shows how the variance is updated.

$$K(t) = \frac{v(t-1)}{v(t-1) + R}, \quad (7)$$

$$y(t) = K[x(t) - y(t-1)] + y(t-1), \quad (8)$$

$$v(t) = v(t-1) - Kv(t-1). \quad (9)$$

The value of  $R$ , the variance in the measurement noise, and the initial value of  $v$ , the variance in the data, were estimated from the training data that were used to train the neural network. The Kalman filter was used to keep a continuously updated estimate of the state of the error curve. The variance  $v$  also suggests bounds on the error of the prediction. These bounds were used to define landmarks — a landmark was placed wherever the actual observation differed from the prediction by more than four standard deviations. The value at which a data point becomes an outlier (here: four



standard deviations) can be altered to select greater or fewer landmarks. The four standard deviation values were chosen through an investigation of the data that were used for training, with the aim being to select landmarks approximately every 2 m.

## 5. Assessing the landmark prediction mechanisms

This work investigates three different neural network architectures (the modified perceptron, the lagged perceptron and the recurrent perceptron) for model acquisition, all with and without median filtered inputs, and two methods for selecting landmarks from the error curves of these networks, peak detection and the Kalman filter.

In order to be able to evaluate the performance of all these different methods it is, therefore, necessary to have some metric so that we can compare them. Two different assessment methods are introduced in this section. Both work by comparing pairs of runs in one environment. Where we have 10 test runs in an environment, we compare the 1st run with the 2nd, 3rd and so on, the 2nd with the 3rd and 4th, etc., so that we obtain 45 pairs for comparison ( $9 + 8 + 7 + \dots + 1 = 45$ ).

The first evaluation method investigates how well two error curves obtained in different runs match, by calculating the sum-squared difference between the two curves. The second evaluates the consistency of the landmark selections, where a landmark is considered to be consistent if it is selected in the same place repeatedly on successive runs.

### 5.1. Assessment by sum-of-squares error

A standard way of comparing two ordered datasets is to measure the square of the distance between the two sets at each point, and to sum these values over the number of elements. This measure increases rapidly when the distance between two points is large, which is a useful feature for our purposes because small deviations between the error curves on two runs are not a problem, but a large difference is likely to lead to a landmark being predicted in one run, but not the other.

Comparing the error curves before the different landmark selection mechanisms are applied allows the performance of the different neural network architectures to be evaluated. It would be expected that

good performance would mean that the error curves would be similar every time.

### 5.2. Assessment by measuring the consistency of landmark selection

The second measure that is proposed is a way of evaluating the consistency of a landmark. A landmark is of high consistency if it is detected in the same place on each run. This means that landmarks that do not correspond to reliable features of the environment, e.g., errors in sensor readings and cross-talk between sensors, are not consistent landmarks. The following measurement, termed LandMark Consistency (LMC) measures this, but allows for some leeway so that a landmark that has moved by one or two sampling points is also given some credit. The measurement is conducted in the following way.

For each test run in a particular environment, a vector is made. There is one element of the vector for each sampling point in the environment. Each element of the vector contains information about whether or not that place has been selected as a landmark. If it is, the value  $1/\sqrt{2}$  is used; if the place is adjacent to a landmark the value  $\frac{1}{2}$  is used, and otherwise a 0 is put into the element. This produces a vector with as many elements as there were samples in the environment, a typical section of which may look like

$$\left( \dots 0, 0, \frac{1}{2}, \frac{1}{\sqrt{2}}, \frac{1}{2}, 0, 0, \dots \right). \quad (10)$$

The vectors for two different runs are then multiplied together using the scalar product, and the average for each landmark is taken by dividing the result by the number of landmarks detected in the run. If the number of landmarks is different in the two runs, the maximum is taken. This means that if the number of landmarks is not the same in both runs, the value of LMC is lower. The values  $1/\sqrt{2}$  and  $\frac{1}{2}$  were chosen so that a perfect match between a landmark in two different runs gives a value of 1, and a close miss gives a value of  $1/\sqrt{2}$ . The information about a landmark being present was spread across three elements of the vector to allow for error in the measurement of the robot's odometry, which could change the position of the landmark. In the event that two consecutive positions were recognised as landmarks, both landmarks would be marked as  $1/\sqrt{2}$ . In this case, a perfect match would result in

a value greater than 1 being produced. This did not happen in any of the testing. The measure can be written in the following way:

$$\text{LMC}(\mathbf{a}, \mathbf{b}) = \frac{\mathbf{a} \cdot \mathbf{b}}{\max(N_a, N_b)}, \quad (11)$$

where  $N_a$  is the number of landmarks in vector  $\mathbf{a}$  (similarly for  $N_b$ ), and LMC the LandMark Consistency performance measure.

Using this measure, each comparison between two runs yields a value between 0 and 1 (assuming that landmarks never appear in consecutive elements), where 0 means that no landmarks correlated, and 1 means that all the landmarks matched perfectly. The two possible different overlaps (of two elements and one element) produce values of  $1/\sqrt{2}$  and  $\frac{1}{4}$ , respectively.

## 6. Results

### 6.1. Data collection

The mechanism proposed here selects landmarks based on the robot's perceptions. The system is entirely passive — it receives data from the sensors, but does not have any control of the motor actions. For this reason, although the system is designed to run

on-line while the robot is in operation, using data collected by the robot in a number of environments is sufficient. On-line operation is demonstrated and discussed in Section 6.5.

The data used in the experiments described in this paper were collected on two different robots. The first was the Bremen wheelchair, *Rolland*, which is shown in Fig. 6. The robot is a motorised wheelchair, made by the German company Meyra, which has been augmented with 27 sonar sensors arranged around the robot. The positions of the sensors are shown in the right of Fig. 6. The wheelchair was used to collect data in seven different environments (A–G) on the 8th floor of the MZH building at the University of Bremen. The layout of this building, and the location of the various data runs are shown in the left of Fig. 8.

The second robot was the Manchester Nomad 200 robot, *FortyTwo*, shown in Fig. 7. The robot has 16 sonar sensors arranged in a ring around its circumference. This robot was run in a number of environments on the second floor of the Computer Science department at Manchester University. A map of this area is given in the right of Fig. 8.

The experimental procedure was the same for all experiments, both in Bremen and Manchester, and consisted of the following steps. The robot was placed at a starting location in the environment. From there the robot moved forwards, pausing every 20 cm and

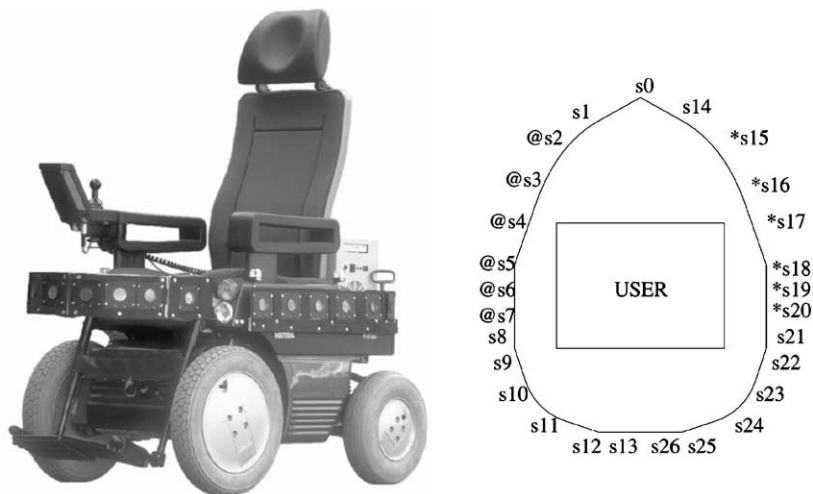


Fig. 6. Left: the Bremen semi-autonomous wheelchair, *Rolland*. Right: the arrangement of the 27 sonars on the wheelchair. @ and \* denote the sonars that were used for data collection.

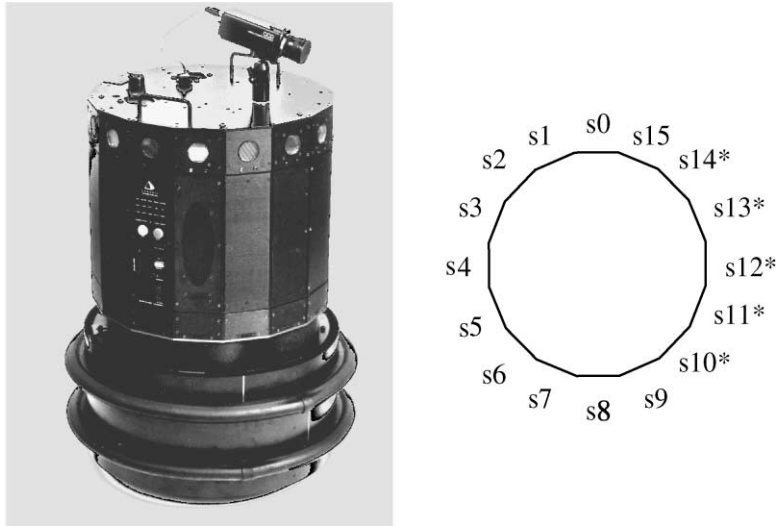


Fig. 7. Left: the Manchester Nomad 200, *FortyTwo*. Right: the arrangement of the 16 sonars on the robot. \* denotes the sonars that were used for data collection.

recording the values of the sonar readings at the current place. The path of the robot through the environment was constrained to be a straight line (or as close as possible) parallel to the wall. The trajectory was controlled by steering with the joystick on the wheelchair, and using a wall-following behaviour produced by a

force-field approach on *FortyTwo*. Further details are given in [7] for the wheelchair and [9] for *FortyTwo*.

The technique of early stopping, described in Section 3.3, was used to decide when network training should be terminated. This was done by obtaining data from 10 runs through an environment, and using five

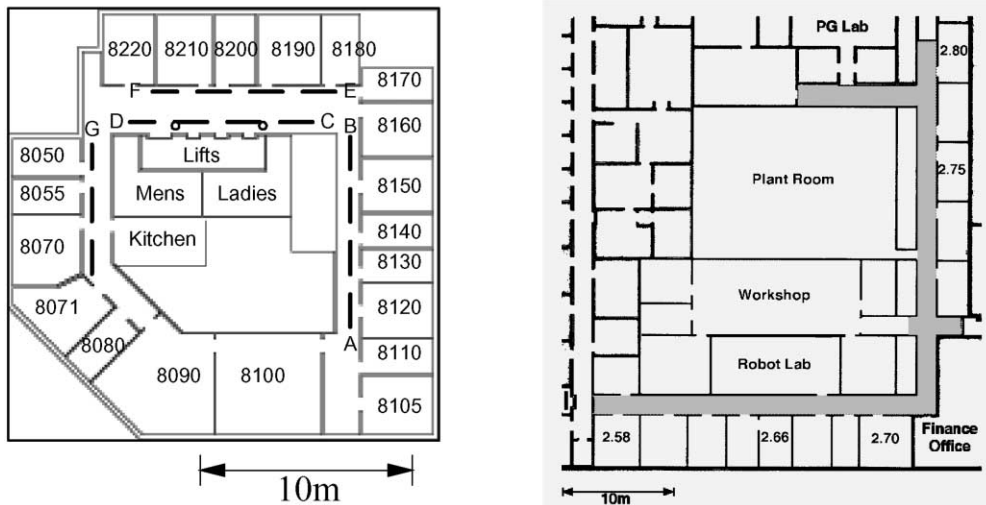


Fig. 8. Left: the environments on the 8th floor of the MZH building at Bremen University where data was collected. The letters show where that data run started, and the dashed lines show the intended robot trajectories. Right: the 2nd floor of the Computer Science Department at the University of Manchester, where further data were collected.

of them as training data and the other five runs as validation data. Using this method, it was established that 20 training epochs was sufficient; the data of those 10 runs was then discarded.

For the experiments proper, data were obtained from another 20 data collection runs in each of a number of environments. Of these 20, 10 were used as training data and 10 were used as test data. The question of how many of the environments should be used to generate the training inputs was investigated, experiment-

ing with taking training data from three environments and from seven environments, the maximum number available. However, much data are used, it should be possible to use the trained network in any environment, including those that it was trained in.

6.2. Evaluation using the sum-of-squares error

Comparing the performance of the trained neural networks before the landmark selection algorithm is

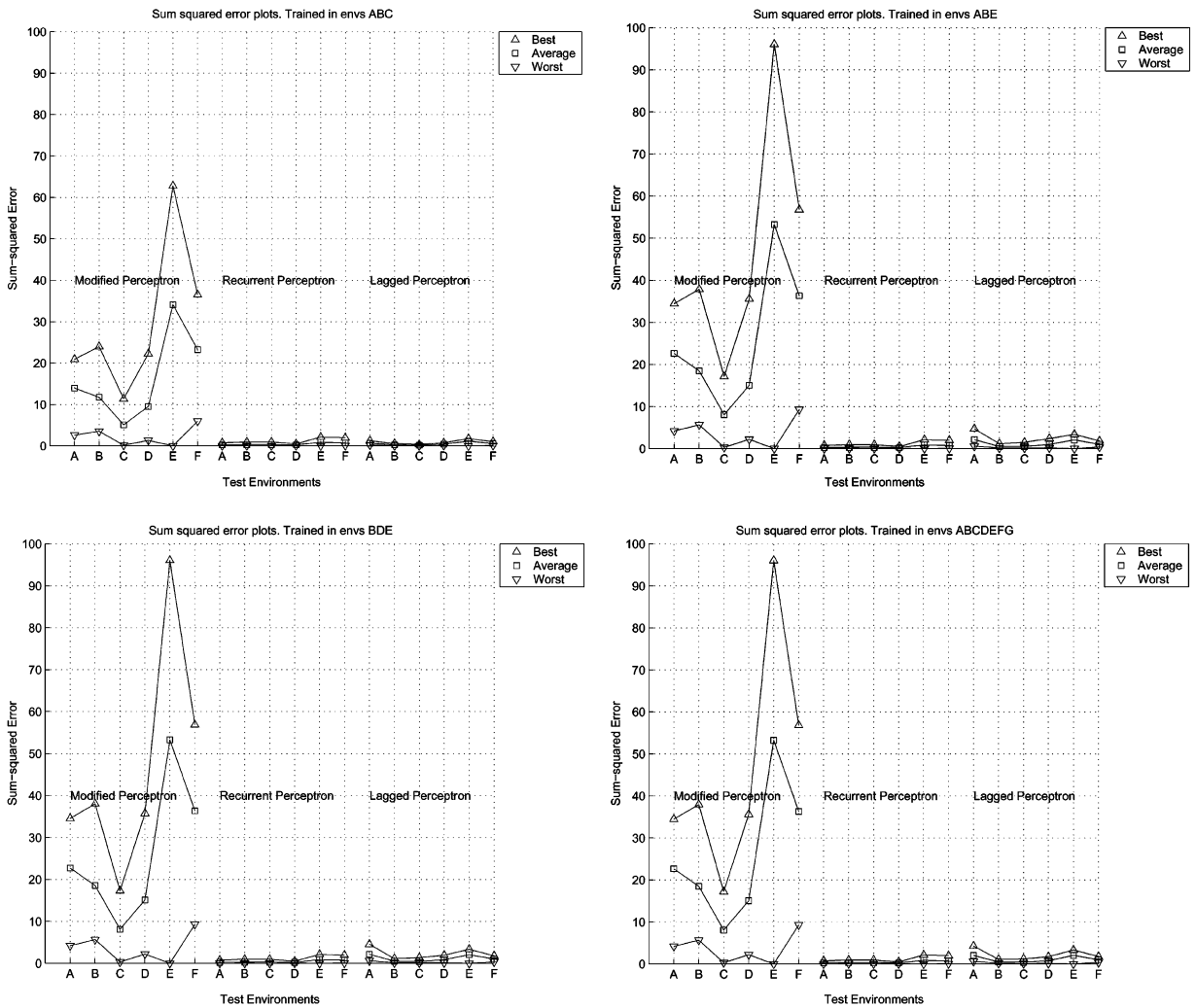


Fig. 9. The results of evaluating the sum-of-squares error for the output of each of the different neural networks. The error is shown for each network, tested in each of environments A–F. The four pictures show the results after training in different sequences of environments (ABC, ABE, BDE, ABCDEFG, respectively).

Table 1

The mean sum-squared error and the standard deviation, for the neural network error curve, averaged over the testing environments, for the different network training algorithms<sup>a</sup>

Algorithm	Training environments	Mean error	Standard deviation
Modified perceptron	ABC	16.28	10.61
	ABE	25.62	16.48
	BDE	25.68	16.45
	ABCDEFG	25.64	16.47
Recurrent perceptron	ABC	0.44	0.28
	ABE	0.64	0.36
	BDE	0.32	0.32
	ABCDEFG	0.45	0.28
Lagged perceptron	ABC	0.53	0.36
	ABE	1.21	0.72
	BDE	1.17	0.75
	ABCDEFG	1.13	0.73

<sup>a</sup> The systems were tested using independent data from training sets collected in environments A–G.

used can give useful information about how consistently the networks are performing, and which features are useful for landmark selection. This comparison is done using the sum-of-squares error. The results given in this section were generated using the data collected on the wheelchair at Bremen University.

The different trained networks were all evaluated over the 10 test runs in each environment. The

Table 2

The mean sum-squared error and the standard deviation, for the neural network error curve, averaged over the testing environments, for the different network training algorithms with the inputs median filtered<sup>a</sup>

Algorithm	Training environments	Mean	Standard deviation
Modified perceptron	ABC	2.69	2.11
	ABCDEFG	2.42	2.32
Recurrent perceptron	ABC	0.51	0.27
	ABCDEFG	0.78	0.66
Lagged perceptron	ABC	0.73	0.67
	ABCDEFG	0.71	0.77

<sup>a</sup> The systems were tested using independent data from training sets collected in environments A–G.

sum-squared-difference between each pair of curves was then evaluated, leading to 45 (9 + 8 + ... + 1) values for each network in each environment. These numbers were averaged to produce an average error for each network in each environment. Fig. 9 shows the average, best, and worst error values for each of the network architectures tried. In addition, Table 1 shows the mean sum-squared error and the standard deviation, averaged across the different test environments. These results show that the modified perceptron generally produces the highest errors. This makes sense since this network receives less input

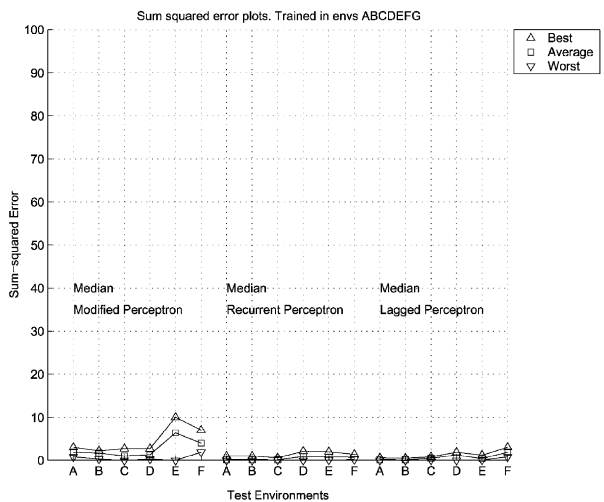
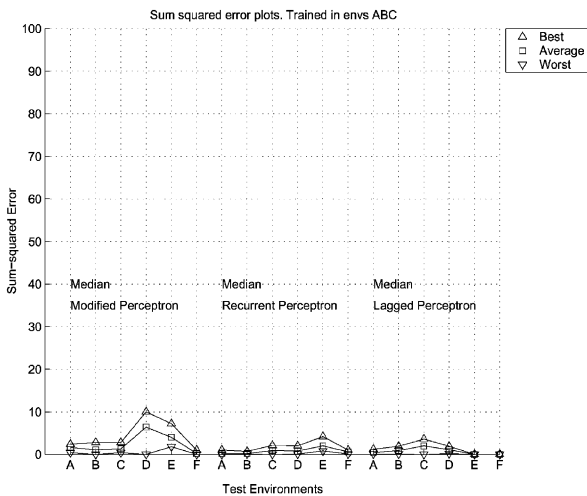


Fig. 10. The results of evaluating the sum-of-squares error for the different neural networks with median filter pre-processing. By comparing with Fig. 9 we can see that it does not make a difference for the lagged and recurrent perceptrons. The error is shown for each network, tested in each of environments A–F. The pictures show the results after training in environments ABC and ABCDEFG, respectively.

that the other networks. The huge errors appear to be caused by places where a landmark is predicted in some runs, but not in others.

By comparing the graph for training in all environments (ABCDEFG) at the bottom right of Fig. 9 with the others, it can be seen that the additional training does not make a significant difference compared to training in only three. A paired Student's *t*-test [10] comparing the sum-squared error after training in environments BDE and in environments ABCDEFG

confirmed this ( $p = 0.01$ ). Using the *t*-test on the sum-squared error curves for the modified perceptron and the lagged perceptron suggests that the chance of them coming from the same distribution is low. The probability of the results of the lagged and recurrent perceptrons being from the same distribution is even lower.

Fig. 10 shows the sum-squared error curves generated when the inputs to the networks are first median filtered, and Table 2 gives the mean and standard

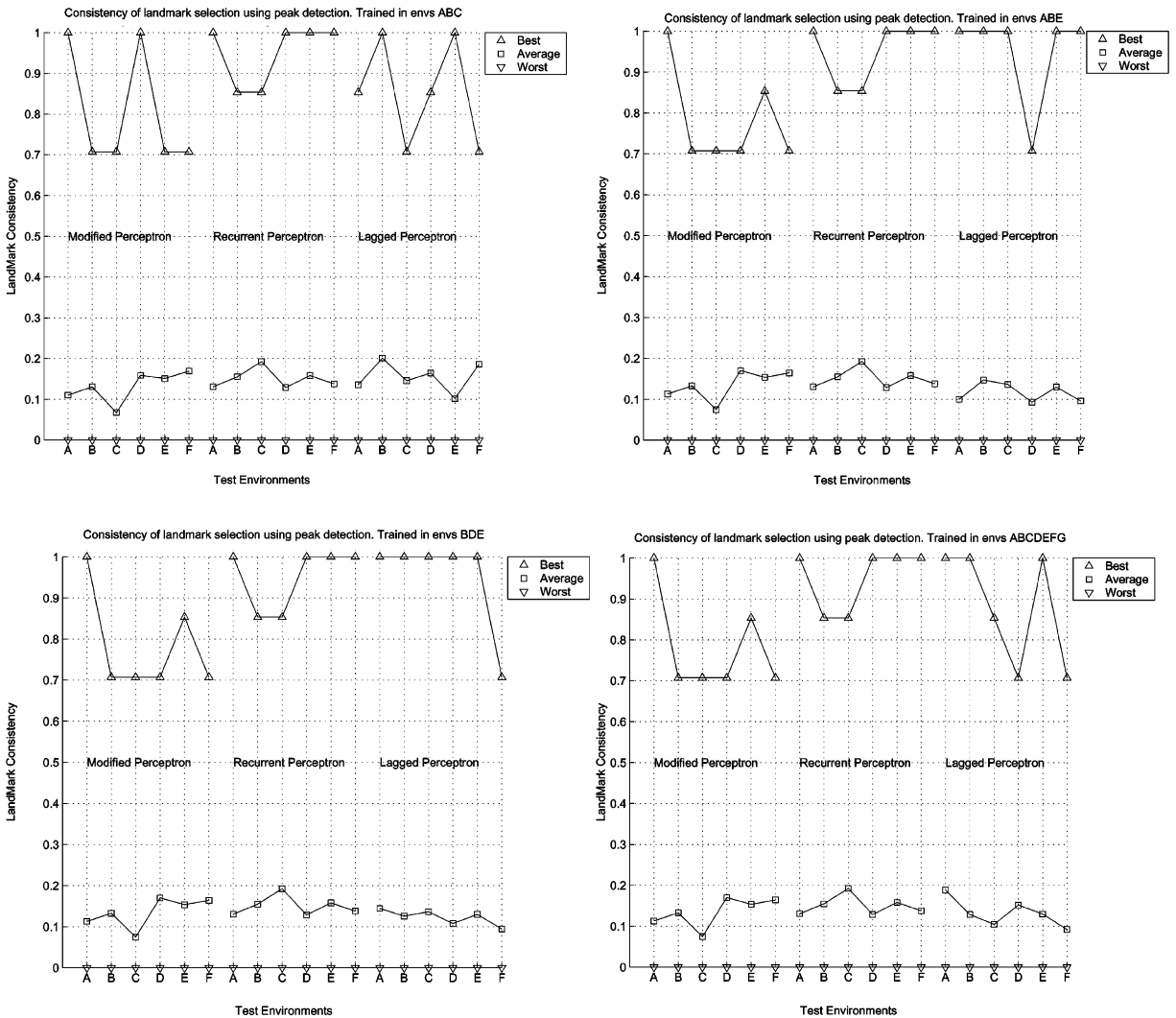


Fig. 11. The results of evaluating the consistency of landmarks (LMC) for the peak detection approach. The LMC is shown for each network, tested in each of environments A–F. The four pictures show the results after training in different sequences of environments (ABC, ABE, BDE, ABCDEFG, respectively).

deviation in the measurements. The results are shown for training in three environments (A–C) and for training in all seven environments. One can see that this technique makes a significant difference to the modified perceptron, but does not really affect the lagged and the recurrent perceptron models. In fact, the results look slightly worse. These results are confirmed by the paired *t*-test, which suggests that the median filtered versions of the recurrent and lagged perceptions correlate highly with the non-median filtered versions, while the chance of the results from

the modified perceptron and the modified perceptron using median filtered input coming from the same distribution is negligible ( $p = 0.01$ ).

### 6.3. Evaluating the landmark consistency

Having compared the quality of the neural network error curves, we now compare the consistency of landmark selection, using the LMC performance measure described in Section 5.2. This was done in a very similar way to the previous sum-squared error measure-

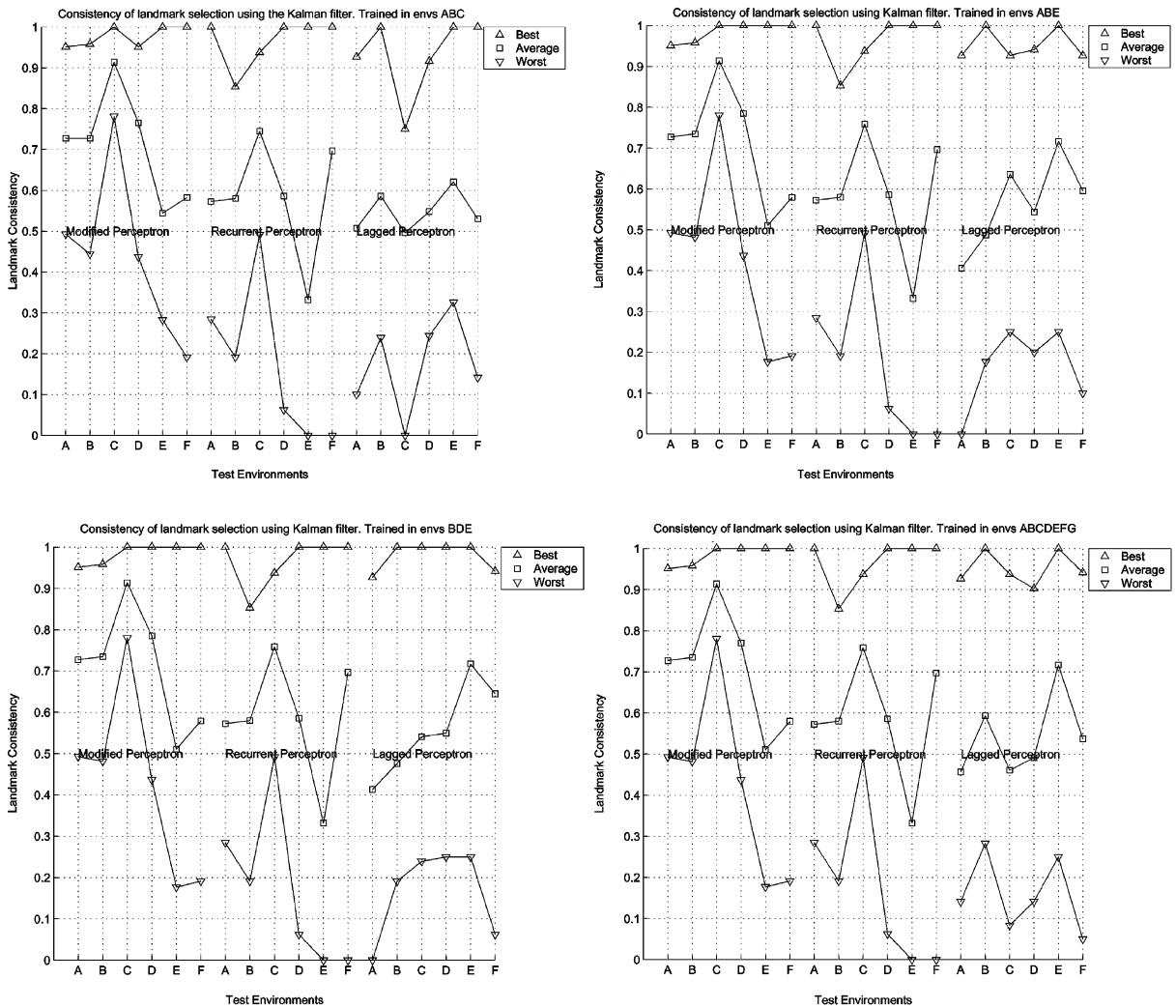


Fig. 12. The results of evaluating the consistency of landmarks (LMC) for the Kalman filter approach. The LMC is shown for each network, tested in each of environments A–F. The four pictures show the results after training in different sequences of environments (ABC, ABE, BDE, ABCDEFG, respectively).

Table 3

The mean LMC and the standard deviation, averaged over the testing environments, for the peak detection and Kalman filter algorithms<sup>a</sup>

Algorithm	Training environments	Peak detection (mean $\pm$ standard deviation)	Kalman filter (mean $\pm$ standard deviation)
Modified perceptron	ABC	0.13 $\pm$ 0.04	0.71 $\pm$ 0.13
	ABE	0.14 $\pm$ 0.04	0.71 $\pm$ 0.15
	BDE	0.17 $\pm$ 0.03	0.71 $\pm$ 0.14
	ABCDEFG	0.15 $\pm$ 0.04	0.71 $\pm$ 0.13
Recurrent perceptron	ABC	0.15 $\pm$ 0.04	0.59 $\pm$ 0.14
	ABE	0.14 $\pm$ 0.05	0.59 $\pm$ 0.15
	BDE	0.15 $\pm$ 0.02	0.61 $\pm$ 0.15
	ABCDEFG	0.15 $\pm$ 0.04	0.61 $\pm$ 0.14
Lagged perceptron	ABC	0.15 $\pm$ 0.02	0.54 $\pm$ 0.05
	ABE	0.14 $\pm$ 0.05	0.56 $\pm$ 0.11
	BDE	0.15 $\pm$ 0.04	0.56 $\pm$ 0.11
	ABCDEFG	0.14 $\pm$ 0.04	0.54 $\pm$ 0.09

<sup>a</sup> The systems were tested using independent data collected in environments A–G.

ments, using an average of the results for each of the testing runs.

The results of this are shown in Figs. 11 and 12 for the three different neural network architectures (modified perceptron, lagged perceptron and recurrent perceptron) with the peak detection algorithm and the Kalman filter, respectively. Again, the average case together with the best and worst cases are shown. Table 3 shows the mean and standard deviation for these results. Again, these are averaged across all of the test environments.

A comparison between the two figures shows very clearly that the performance of the Kalman filter with this measure is significantly better than that of the peak detection. The paired Student's *t*-test was used to compare the peak detection and Kalman filter algorithms for each of the neural network architectures. The results show that the probability of the LMC measures for the two algorithms coming from the same distribution, regardless of neural network used, is negligible.

Looking at Fig. 11, which shows the LMC measure for the peak detection, we see that while the best case performance is fairly good, in the worst case there are no matches, and the average is also very low. The performance of the algorithm does not appear to vary much between the different network architectures, nor between the different training sets. A paired *t*-test comparing the different network architectures shows that the performances of the lagged perceptron and the modified perceptron are very similar ( $p = 0.01$ ),

while the performance of the recurrent perceptron is significantly different (in both cases,  $p = 0.01$ ).

Fig. 12 shows the results of the LMC evaluation for the Kalman filter algorithm. It can be seen that the average landmark consistency is significantly higher than it is for the peak detection, and the worst case has risen above 0 in virtually every case. Interestingly, the Kalman filter technique appears to work best on the modified perceptron data, which the sum-of-squared error method of the previous section found to be most noisy. This combination of modified perceptron and Kalman filter provides the best landmark consistency overall, which leads to the question of whether or not we actually need the neural network, rather than

Table 4

The average number of landmarks chosen in each environment by the two landmark detection algorithms<sup>a</sup>

The environment	Average number of landmarks $\pm$ standard deviation	
	Peak detection	Kalman filter
A	2.1 $\pm$ 0.5	4.5 $\pm$ 1.1
B	2.5 $\pm$ 1.0	5.5 $\pm$ 0.5
C	1.8 $\pm$ 0.7	3.4 $\pm$ 0.9
D	1.0 $\pm$ 0.0	5.0 $\pm$ 0.0
E	1.4 $\pm$ 0.5	4.9 $\pm$ 0.5
F	1.9 $\pm$ 0.6	3.4 $\pm$ 0.6

<sup>a</sup> The neural network that was used was the one that performed best for each of the landmark algorithms, the recurrent perceptron for the peak detection and the modified perceptron for the Kalman filter.



a very simple model that just predicts that the new perception is the same as the current one. However, while simpler, this system has a number of disadvantages. For instance, whenever a doorway is predicted there will be a large number of potential landmarks detected, as the various sensors detect the doorjamb, followed by the door itself and then the second doorjamb.

#### 6.4. Number of landmarks

Table 4 shows the average number of landmarks that are produced by each of the algorithms for different test sets. It can be seen that the Kalman filter approach finds significantly more landmarks, and that the standard deviation in the number of landmarks found is similar for both landmark detection algo-

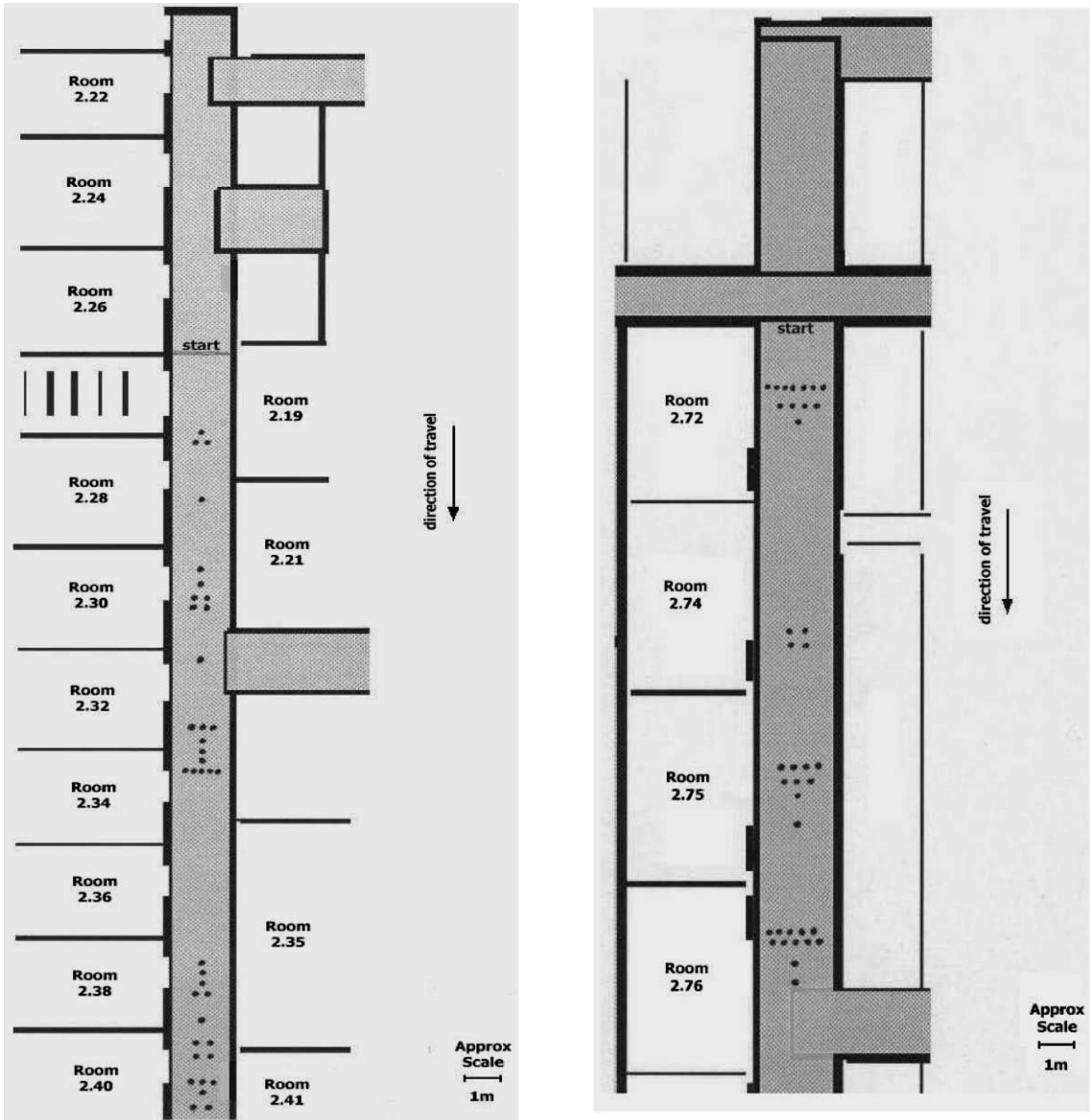


Fig. 13. The positions of the landmarks (drawn as dots) using the peak detection algorithm on two different environments.

rithms, averaged over the different neural network architectures. The goal of this project is to reduce the number of landmarks that are needed for reliable robot navigation, but obviously there still need to be a usable number of landmarks. As the results given here show that in general the landmarks are not found perfectly, having several landmarks in an environment is a definite advantage, as it increases the probability of the robot finding some of them.

### 6.5. On-line operation

In addition to the off-line operation that has been discussed in previous sections, we also want to demonstrate that the algorithms can be used on-line. Once the neural network has been trained, the processing of the sonar data is a very simple task and does not require much computational time. It is, therefore, suitable for on-line use. A version of the algorithms that detected landmarks for use as the robot explored was implemented.

Using the peak detection algorithm operating on *FortyTwo*, the robot travelled along a number of different corridor environments. Each time the algorithm selected a landmark, the robot notified the user, who noted the position of the landmark on a floor plan. The robot travelled in each environment 12 times. The results of two of these experiments are shown in Fig. 13. The picture on the left demonstrates one reliable landmark, picked in 10 of the 12 runs, outside room 2.32, and one that is spread across several sampling locations (outside room 2.30). The area outside room 2.38 appears as one large landmark.

The results obtained in a second environment, shown in the right of Fig. 13 are crisper. There are several stretches of corridor where no landmarks are detected in any of the runs. These mark places where the perceptions of the robot are predicted accurately by the network, so that no landmarks are selected. We can also see three good clusters of landmarks, where a landmark was predicted reliably, and only one area where the prediction was less reliable.

## 7. Conclusions

### 7.1. Motivation

For a mobile agent, the ability to navigate is of fundamental importance. Over short distances, this

capability of goal-oriented motion may be achieved by path integration through dead reckoning, but very quickly odometry drift errors render dead reckoning navigation systems too inaccurate for useful navigation.

The alternative is to use perceptual landmarks for navigation. Typical landmark-based navigation systems either log perceptions at regular intervals, or match perceptions with pre-supplied templates of ‘desirable’ landmarks. The former approach is inefficient and runs the risk of missing out clearly visible landmarks that lie between sampling points. The latter approach runs the risk of being based on landmarks that are easily visible to a human, but not to a robot, or (equally undesirable) of missing landmarks that are easily visible to a robot, but that remain unselected because they are not important for human navigation.

A suitable way to address these problems is to devise an automatic landmark selection mechanism that allows the robot to select suitable landmarks without a pre-defined model of those landmarks, or human supervision. This paper has described an algorithm that can be used to achieve this.

### 7.2. Approach

In this paper, it is argued that a landmark should be both clearly detectable and conspicuous, and reliably and repeatedly detectable. To identify conspicuous perceptions a robot acquires a model of the usual temporal relationship between consecutive sensory perceptions. In the landmark-selection phase, this model is then used to predict the next sensory perception, based on the current sensory perception. Wherever this prediction fails, i.e., wherever the predicted perception differs markedly from that actually observed, the location is selected as a landmark on the grounds that it ‘stands out’ from the usual perceptions commonly observed. To assess the consistency of landmark selection, we use two performance measures — sum-squared error and LMC.

### 7.3. Discussion of results

Three different neural networks were investigated as ways to acquire the model, and two different methods of selecting landmarks based on the prediction error

made by the respective model were tried. Experiments show that landmark selection is possible and consistent using the proposed method, and that this research is promising enough to merit further investigations on an even larger scale. The three models investigated — modified perceptron, recurrent perceptron and lagged perceptron — do not differ dramatically in their prediction accuracy.

When selecting landmarks by detecting peaks in the error curve, the performance of the modified perceptron is worst, while the recurrent perceptron and lagged perceptron perform practically identically (Fig. 9). However, with appropriate pre-processing of the raw sensor perceptions, the modified perceptron also performs as well as the other two models (Fig. 10).

If the Kalman filter is used for landmark selection, there is no discernible difference in performance between the three models (Fig. 12), but the results are always better than using peak detection. The Kalman filter approach has another possible benefit — because the definition of an ‘outlier’ (currently four standard deviations) can be modified, the number of landmarks selected in a particular environment can also be influenced. This property is useful, because it facilitates the application of more stringent selection criteria in ‘landmark-rich’ environments, and less stringent criteria in ‘landmark-poor’ environments. However, the experiments reported here show that a value of four standard deviations produces useful results in a variety of environments.

The results also show that in order to train the model it is not necessary to expose it to *all* environments that the robot may encounter during operation (Fig. 9). This observation confirms initial expectations. The robot is essentially learning a model of the temporal relationship between its own sensory perceptions, not a model of any particular world. Provided that the training data are rich enough to provide most possible temporal constellations of sensory perceptions, the model acquired in one or two training environments is then applicable to *any* environment. As the training data are acquired during several runs through an environment, the robot will have taken paths that have slightly different orientations. It can, therefore, be assumed that these small changes will be built into the model.

There is a further interesting application of the method described here. Obviously, the robot acquires

a model of ‘normal’ perceptions against which each set of sensory perceptions are measured, to detect those perceptions that ‘stand out’. This is a form of novelty detection. The algorithm described here can, therefore, be used to detect abnormalities (with respect to the robot’s sensory perceptions) in the environment.

#### 7.4. Future work

To make predictions, the robot has to move in straight-line segments between consecutive perceptions. It is conceivable that the approach can be expanded to arbitrary motions between perceptions. Furthermore, for the experiments described here, the robots have sampled perceptual data in regular intervals. Clearly, the proposed algorithm is intended to select landmarks from *continuous* observations of the environment and although a robot will always only be able to sample the environment in discrete time steps, it is no problem to increase the sampling rate beyond that used in the experiments discussed in this paper.

Finally, how useful the selected landmarks are for actual navigation has not yet been investigated. The purpose of this research was to investigate the *consistency* of landmark selection. The construction of an entire mobile robot navigation system, incorporating the landmark selection algorithm presented here, is subject to ongoing research.

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