Prevalence, clinical features and risk assessment of pre-diabetes in Spain: the prospective Mollerussa cohort study

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ABSTRACT

Purpose The Mollerussa prospective cohort was created to study pre-diabetes in a population-based sample from the primary care setting in the semirural area of Pla d’Urgell in Catalonia (Spain). The aims of the study were to assess the prevalence of pre-diabetes in our population, the likelihood to develop overt diabetes over time and to identify risk factors associated with the progression of the condition.

Participants The cohort includes 594 subjects randomly selected between March 2011 and July 2014 from our primary care population, who were older than 25 years, consented to participate and did not have a recorded diagnosis of diabetes.

Findings to date At baseline, we performed a clinical interview to collect demographic, clinical and lifestyle (including a nutritional survey) characteristics; carotid ultrasound imaging to assess subclinical cardiovascular disease was also performed, and a blood sample was collected, with an overall <5% rate of missing data. An additional blood draw was performed 12 months after initial recruitment to reassess laboratory results in patients initially identified as having pre-diabetes, with an 89.6% retention rate. Several studies investigating various hypotheses are currently ongoing.

Future plans All subjects recruited during the cohort creation will be followed long-term through annual extraction of data from health records stored in the electronic Clinical station in Primary Care database. The Mollerussa cohort will thus be a sound population-based sample for multiple future research projects to generate insights into the epidemiology and natural history of pre-diabetes in Spain.

INTRODUCTION

According to the American Diabetes Association (ADA), diabetes is broadly classified into four categories: type 1 diabetes mellitus (T1DM), type 2 diabetes mellitus (T2DM), gestational diabetes and specific types of diabetes due to other causes. However, there is a group of individuals that, in spite of having higher than normal glucose levels, do not meet criteria for diabetes, a condition referred to as pre-diabetes.

There are different definitions of pre-diabetes, but the most common one, the ADA criteria, considers one of the following instances: (a) impaired fasting plasma glucose (IFG), defined as fasting plasma glucose (FPG) between 100 and 125 mg/dL (5.6–5.9 mmol/L); or (b) impaired glucose tolerance (IGT), defined as a 2-hour plasma glucose value after a 75g oral glucose tolerance test (OGTT) between 140 and 199mg/dL (7.8–11.0 mmol/L); or (c) glycated haemoglobin (HbA1c) levels between 5.7% and 6.4% (39–46mmol/mol).

The prevalence of pre-diabetes varies across countries and depending on the parameter used for the estimations. Based solely on IGT, its worldwide prevalence among adults has been estimated by the International Diabetes...
Federation to be 6.7% in 2015, with half of them (50.1%) younger than 50 years. In England, solely based on HbA1c levels, the prevalence was 35.5% among the adult population in 2011; in Spain, isolated IFG and isolated IGT were present in 3.4% and 2.9%, respectively, and combined IFG–IGT in 2.2% of the adult population in 2010; and in the USA, using the ADA definition (HbA1c levels or IFG or IGT) the prevalence was as high as 38% in 2012.

Understanding the epidemiology and natural history of pre-diabetes has become a health priority, in particular at the primary care setting, because it is a source of avoidable morbidity and mortality. First, individuals with IFG and/or IGT have a clinical phenotype that resembles patients with T2DM, as they tend to be older, have a higher body mass index (BMI), have more frequent insulin resistance and dyslipidaemia and have higher arterial blood pressure (BP) than people with normal glucose tolerance. Second, people with pre-diabetes are at increased risk of developing diabetes: according to the ADA, up to 70% of them will eventually develop overt diabetes; the annual incidence of progression to diabetes is around 5%–10% depending on the population characteristics and the definition of pre-diabetes; 6%–9% in subjects with isolated IFG, 4%–6% in those with isolated IGT, up to 15%–19% among those with both IFG and IGT and subjects with HbA1c levels from 5.7% to 6.5% have a 7.5-year predicted risk of 43.1% for incident diabetes. Finally, individuals with pre-diabetes are at increased risk of cardiovascular disease (CVD) and premature mortality: a meta-analysis found that the risk of CVD is increased regardless of type of blood glucose assessment in comparison to subjects with normoglycaemia; and another recent meta-analysis found that risks of all-cause and CVD mortality compared with normoglycaemic subjects were increased in individuals with pre-diabetes with ADA defined IFG, IGT or both criteria combined, although not with isolated IFG.

Based on epidemiological and clinical evidence, it is important from a prediction and prevention perspective to target segments of the population with metabolic and cardiovascular (CV) signatures associated with an increased risk of developing diabetes and CVD. The Mollerussa cohort was designed to identify undiagnosed diabetes or pre-diabetes in the adult primary care population from a semi-rural area of Catalonia (Spain), and to further obtain extensive epidemiological, clinical (including subclinical atherosclerotic disease) and lifestyle data. In the following phases, the cohort will be run as prospective observational studies involving identified at-risk individuals to determine the progress over time regarding risk factors, incident diabetes, incidence of CV events, rates of hospitalisation and global mortality.

**COHORT DESCRIPTION**

Mollerussa is a prospective observational cohort study conducted in Pla d’Urgell, a semi-rural area of Catalonia (Spain), among subjects from the general population with healthcare coverage from the Catalan Institute of Health (Institut Català de la Salut).

Based on an estimated prevalence of pre-diabetes in the area of Lleida of 11% (10%–19%) in 2011, we initially calculated that we would need a representative sample of 940 adults considering a 95% CI and a margin of error of ±2%. However, literature published after the initiation of the recruitment phase, using HbA1c levels and ADA criteria, reported pre-diabetes prevalence between 35.5% and 38%. Using this datum, a random sample of 505 subjects was sufficient to assess an estimated prevalence of about 30% with a 95% CI and a margin of error of ±4%.

**Recruitment**

Using the electronic Clinical station in Primary Care (eCAP) health records database implemented in all primary care centres in Catalonia, a code number was given to each registered adult. During the recruitment phase, 24,666 registered health records met inclusion criteria, namely subjects older than 25 years and attending any Primary Healthcare Centre in the same health area in Pla d’Urgell (box 1; figure 1); among them, 2,226 random individuals (about 5% of the total number of individuals registered at each centre) were contacted by telephone (up to three attempts) and invited to participate (figure 1). Randomisation was carried out using a randomiser programme (SPSS software V.16.0 for Windows; SPSS), following the principles of simple random sampling.

Main exclusion criteria (box 1) included a previous diagnosis of diabetes (T1DM, T2DM and any specific subtype of diabetes), and subjects on treatment with oral antidiabetic drugs to avoid the inclusion of individuals with actual diabetes but optimal glycaemic control, or even without diabetes but using metformin as treatment for other conditions. Based on their willingness to join the study and exclusion criteria, a total of 602 subjects were recruited and subsequently invited for an appointment, of whom four withdrew consent and, for four, we did not obtain any baseline laboratory data, therefore giving a final study population of 594 subjects.

**Data collection**

The research plan included a first phase involving two visits to the primary healthcare centre for baseline screening, a second phase conducted 12 months after the baseline visits and a third phase of long-term follow-up of the cohort.

**First phase or baseline screening**

The following variables were collected/explored by trained research staff in the first visit to the subject’s primary healthcare centre: (a) Sociodemographic variables: age; gender; education level according to the International Standard Classification of Education; sector of working activity (primary, secondary or manufacturing, tertiary or services); self-perceived work activity (minimum, light, moderate, heavy); report on...
Inclusion criteria
Age ≥25 years
Attended a primary health care centre in the area

Exclusion criteria
Patient information about having diabetes provided in the first contact or existing ICD-10 code of diabetes (E11, E14 or E13) registered by a physician or confirmed based on clinical data:
- HbA1c ≥6.5%
- IGT: 2-hour plasma glucose in the 75g OGTT ≥200 mg/dL (11.1 mmol/L)
- IFG: FPG ≥126 mg/dL (7 mmol/L)
Specific subtypes of diabetes other than T1DM and T2DM:
- Gestational diabetes
- Genetic defect of beta-cell action
- Genetic defect in insulin action
- Diseases of the exocrine pancreas (eg, pancreatitis, haemochromatosis, pancreatic cancer, cystic fibrosis)
- Endocrinopathies (eg, Cushing’s syndrome, glucagonoma, somatostatinoma, hyperthyroidism, pheochromocytoma, acromegaly)
- Chemical-induced diabetes
- Diabetes secondary to infections
- Autoimmune diabetes
Use of oral antidiabetic drugs: metformin, dipeptidyl peptidase-4 inhibitors, sulfonylureas and glitazones
Presence of cardiovascular disease:
- Previous hospitalisation to treat heart disease
- Heart failure
- Left bundle branch block or second degree atrioventricular block
- Aortic stenosis
- Systolic BP >160 mm Hg or diastolic BP >105 mm Hg
Cancer treated in the preceding 5 years, except non-melanoma skin cancers (basal-cell and squamous-cell carcinoma)
Kidney disease, defined as plasma creatinine ≥1.4 mg/dL in men and ≥1.3 mg/dL in women or proteinuria >2+
Anaemia, defined as haematocrit <36% in men and <33% in women
Hepatitis, defined as transaminases more than 10 times the upper limit of normal
Gastrointestinal diseases (pancreatitis, irritable bowel disease and inflammatory bowel disease)
Recent abdominal surgery
Chronic pulmonary obstructive disease requiring domiciliary oxygen therapy
Chronic infectious diseases (eg, HIV, active tuberculosis, HBV and HCV)
Use of systemic glucocorticoids or beta blockers
Major psychiatric disorder with psychotic symptoms

Box 1 Summary of inclusion and exclusion criteria

In summary, the prevalence of pre-diabetes and diabetes was assessed in a sample of 1586 participants from a primary care setting in Catalonia, Spain. The study found that the prevalence of pre-diabetes was 34.7%, while the prevalence of T2DM was 5.3%. The results suggest that primary care providers should be vigilant in screening for diabetes and pre-diabetes, as these conditions are highly prevalent in the population studied.

Second phase or short-term follow-up
Twelve months after the baseline visits, subjects initially fulfilling pre-diabetes criterion underwent a second visit to perform another blood draw to reassess the laboratory results. This was based on the ADA recommendation to repeat testing in the absence of unequivocal hyperglycaemia. Based on re-evaluated HbA1c levels, those subjects with HbA1c levels between 5.7% and 6.4% were confirmed as pre-diabetes; those with a subsequent increase from pre-diabetes values at baseline to HbA1c ≥6.5% after 12 months were considered as incident diabetes (and as well re-contacted and advised to visit a general practitioner); and those with a further decrease
Figure 1  Mollerussa study flow chart. CH, clinical history; HbA1c, glycated haemoglobin.
from pre-diabetes values at baseline to HbA1c <5.7% after 12 months as regression to normoglycaemia, although they will be followed up to rule out a temporary improvement, and thus a false case of regression to normal mean HbA1c values.

The 12-month follow-up will give a real estimation of pre-diabetes prevalence, valuable information on the 1-year probability of progression to diabetes and which risk factors were relevant to the further development of the disease (eg, metabolic traits and lifestyle).

Third phase or long-term follow-up
In addition to the 12-month follow-up, we also plan to follow all subjects enrolled in the Mollerussa cohort annually through cross-sectional extraction of data stored in the primary care electronic medical records of the eCAP database. This is based on the ADA recommendation to follow pre-diabetes in asymptomatic adults yearly. We will extract data for the following variables: any diagnosis of T2DM (International Classification of Disease (ICD-10) codes E11 or E14); time since diagnosis; estimated glomerular filtration rate using the Modified Diet in Renal Disease formula; standardised HbA1c values, using the most recent value of the preceding 12 months; presence of CVD, including coronary artery disease (ICD-10 codes I20, I21, I22, I23 or I24), stroke (ICD-10 codes I63, I64, G45 or G46) and peripheral artery disease (ICD-10 code I73.9); risk factors, including BMI (most recent value in the last 12 months), cholesterol levels (total, low-density lipoproteins or LDL-cholesterol and high-density lipoproteins or HDL-cholesterol; most recent value in the last 12 months), BP (systolic and diastolic mean value in the last 12 months); and data on prescribed glucose-lowering, lipid-lowering, antihypertensive and antithrombotic medications. This information will be supplemented with data registered in the Conjunto Mínimo Básico de Datos de Altas Hospitalarias (Set of Minimum Basic Data Set of Hospital Admissions), which records all admissions to public and private hospitals in the region and contains information on diagnostics, procedures and discharge reports.

With this additional longitudinal approach, we will be able to obtain the patient’s data on progression to overt diabetes and/or initiation of antidiabetic treatment over time (if directly related to diabetes). This is important because besides the annualised incidence rate of progression to diabetes, the time course progression of dysglycaemia has not been studied at large. From the few available studies, the mode of onset of diabetes in subjects with pre-diabetes follows a non-linear pattern, with a rapid rather than gradual onset of diabetes over a 3-year time. Moreover, we will be able to obtain data on the incidence of other diabetes-associated chronic conditions also present at the pre-diabetes stages, such as nephropathy, neuropathy or retinopathy; the incidence of macrovascular complications over time; the likelihood of initiation of hypoglycaemic agents among progressors; rates and cause of hospitalisations; and overall mortality.

This study was approved by the Ethics Committee of the Primary Healthcare University Research Institute (Institut d’Investigació en Atenció Primària, IDIAP Jordi Gol (P12/043), and all patients signed a written informed consent form prior to participation. The study was conducted in accordance with the Declaration of Helsinki (1964).

**FINDINGS TO DATE**

The Mollerussa study completed its recruitment phase between August 2011 and July 2014, and the 12-month short-term follow-up in July 2015. The enlisted sample (n=602) was different from the eligible but not enlisted population (n=784) in terms of gender and age (table 1); the enlisted sample had a significantly higher proportion of women (58.6% vs 44.0%; p<0.001), and was older (mean age 48.1 years vs 45.7 years; p<0.001).

Demographic characteristics of participants who provided a blood sample during the first phase (baseline visit) are summarised in table 2.

After the first phase, the rate of missing data was less than 5% across variables of interest (table 3), which is far below the 20% maximum recommended lost to follow-up.

### Table 1 Age and gender characteristics between subjects eligible but not enlisted and subjects eventually enlisted in the Mollerussa cohort

<table>
<thead>
<tr>
<th>Age group, years, n (%)</th>
<th>Failed to be contacted (n=448)</th>
<th>Declined to participate (n=349)</th>
<th>All (n=797)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender (female), n (%)</td>
<td>181 (40.4)</td>
<td>170 (48.7)</td>
<td>351 (44.0)</td>
</tr>
<tr>
<td>Age, years, mean (SD)</td>
<td>44 (15.0)</td>
<td>47.9 (16.3)</td>
<td>45.7 (15.7)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>48.1 (13.4)</td>
</tr>
<tr>
<td>Age group, years, n (%)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;40</td>
<td>218 (48.7)</td>
<td>140 (40.1)</td>
<td>358 (44.9)</td>
</tr>
<tr>
<td>40–60</td>
<td>160 (35.7)</td>
<td>125 (35.8)</td>
<td>285 (35.8)</td>
</tr>
<tr>
<td>&gt;60</td>
<td>70 (15.6)</td>
<td>84 (24.1)</td>
<td>154 (19.3)</td>
</tr>
</tbody>
</table>

*χ² test, p<0.001.
†Kruskal-Wallis non-parametric test, p<0.001.

at baseline (n=193 excluding undiagnosed diabetes), a 89.6% of subjects found to have altered glycaemia levels.

During the first and second phase of the Mollerussa cohort pre-diabetes in high-risk individuals. Conversely, HbA1c measurement is cost-effective and improves the sensitivity of FPG in the detection of early T2DM in high-risk individuals.

Table 2 Demographic characteristics of study population enrolled in the Mollerussa cohort

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Total valid N</th>
<th>Statistic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender, women, n (%)</td>
<td>594</td>
<td>347 (58.4)</td>
</tr>
<tr>
<td>Age, years, mean (SD)</td>
<td>594</td>
<td>50.6 (13.3)</td>
</tr>
<tr>
<td>Weight, kg, mean (SD)</td>
<td>574</td>
<td>73.1 (14.5)</td>
</tr>
<tr>
<td>Waist, cm, mean (SD)</td>
<td>573</td>
<td>94.2 (12.5)</td>
</tr>
<tr>
<td>Body mass index, kg/m², mean (SD)</td>
<td>573</td>
<td>26.3 (4.7)</td>
</tr>
<tr>
<td>&lt;25.0, n (%)</td>
<td>236</td>
<td>41.2 (SD)</td>
</tr>
<tr>
<td>25.0–29.9, n (%)</td>
<td>235</td>
<td>41.0 (SD)</td>
</tr>
<tr>
<td>≥30.0, n (%)</td>
<td>102</td>
<td>17.8 (SD)</td>
</tr>
<tr>
<td>Education level, n (%)</td>
<td>575</td>
<td>24 (4.2)</td>
</tr>
<tr>
<td>Not even primary school</td>
<td></td>
<td>393 (68.7)</td>
</tr>
<tr>
<td>Completed primary school</td>
<td></td>
<td>122 (12.2)</td>
</tr>
<tr>
<td>Secondary/high school</td>
<td></td>
<td>366 (63.7)</td>
</tr>
<tr>
<td>Graduate or higher</td>
<td></td>
<td>63 (11)</td>
</tr>
<tr>
<td>Work activity, n (%)</td>
<td>572</td>
<td>393 (68.7)</td>
</tr>
<tr>
<td>Employed</td>
<td></td>
<td>12 (2.1)</td>
</tr>
<tr>
<td>Unemployed</td>
<td></td>
<td>65 (11.4)</td>
</tr>
<tr>
<td>Disability</td>
<td></td>
<td>12 (2.1)</td>
</tr>
<tr>
<td>Retired</td>
<td></td>
<td>102 (17.8)</td>
</tr>
<tr>
<td>Hypertension, n (%)</td>
<td>571</td>
<td>102 (17.9)</td>
</tr>
<tr>
<td>Dyslipidaemia, n (%)</td>
<td>571</td>
<td>131 (22.9)</td>
</tr>
<tr>
<td>Hypertiglyceridaemia, n (%)</td>
<td>554</td>
<td>22 (4.0)</td>
</tr>
<tr>
<td>Smokers, n (%)</td>
<td>594</td>
<td>152 (25.6)</td>
</tr>
<tr>
<td>Current</td>
<td></td>
<td>148 (24.9)</td>
</tr>
<tr>
<td>Former</td>
<td></td>
<td>152 (25.6)</td>
</tr>
<tr>
<td>Alcohol consumption, n (%)</td>
<td>573</td>
<td>286 (49.9)</td>
</tr>
</tbody>
</table>

We are currently in the phase of longitudinal follow-up of all subjects with subjects identified during the first and second phase (figure 1), and developing protocols for the analyses to explore hypotheses on different features of the epidemiology and natural history of pre-diabetes in our primary care setting. The first paper from the Mollerussa project (now in preparation) will describe results derived from the short-term follow-up of the cohort, namely the prevalence of pre-diabetes and undiagnosed diabetes, the clinical and demographic profile of patients with pre-diabetes versus those with normal glycaemic levels, a first estimation of the annual incidence of overt diabetes among subjects with pre-diabetes and the metabolic, CV and lifestyle disease-associated conditions.

STRENGTHS AND LIMITATIONS

The main strength of the Mollerussa cohort is that it includes adult patients from primary care health centres prospectively collected, and the opportunity to follow participants in the long term through healthcare electronic registries, which ensures that it will be of use for multiple future research projects. The combined short-term prospective and long-term longitudinal design has both advantages and limitations.

The prospective phase (baseline screening and 12-month short-term follow-up) prevents recall bias because the risk for diabetes was assessed before the onset of the disease, and the measurement of events in a temporal sequence allows for causes to be distinguished from effects. However, we must acknowledge a potential selection bias, since we had higher rates of women and middle-aged subjects among enlisted people than among eligible but not enlisted subjects. The influence of this potential bias will be minimised through a weighting process on the prevalence estimates, although how this original unbalance may impact the results is not clear, because IFG and HbA1c detect different categories of individuals as being at risk: IFG is substantially more common among men, and its prevalence tends to plateau in middle age, while the prevalence of pre-diabetes using HbA1c increases with age (maximum peak in those aged 60–74 years) but does not differ by gender. Finally, we did not perform an OGTT among enrolled individuals. Although IGT is more common than IFG in most populations, it is more sensitive but slightly less specific for identifying people who will develop diabetes. Additionally, the OGTT has low reproducibility and it is inconvenient in terms of costs and time consumption. Conversely, HbA1c measurement is cost-effective and improves the sensitivity of FPG in the detection of early T2DM in high-risk individuals.

On the other hand, the longitudinal, long-term phase has the advantage that cohort membership is not dependent on continuing to visit the practice from which the members were recruited. While the main strength is that retention rate also indicating acceptable validity of the results (table 3; figure 1).

Table 3 Summary of missing data for variables recorded during the first and second phase of the Mollerussa cohort

<table>
<thead>
<tr>
<th>Variable of interest</th>
<th>Missing data, n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>First phase (baseline screening); n=594</td>
<td></td>
</tr>
<tr>
<td>No clinical interview</td>
<td>17 (2.9)</td>
</tr>
<tr>
<td>No nutritional survey</td>
<td>28 (4.7)</td>
</tr>
<tr>
<td>No carotid echography</td>
<td>17 (2.9)</td>
</tr>
<tr>
<td>No laboratory results</td>
<td>0 (0.0)</td>
</tr>
<tr>
<td>No HbA1c measurement</td>
<td>10 (1.7)</td>
</tr>
<tr>
<td>No sample for biobank</td>
<td>22 (3.7)</td>
</tr>
<tr>
<td>Second phase (12 months follow-up); n=193</td>
<td></td>
</tr>
<tr>
<td>No laboratory results</td>
<td>20 (10.4)</td>
</tr>
</tbody>
</table>

HbA1c, glycated haemoglobin.
this will minimise losses to follow-up, the primary limitation is that it will rely on data that may be incomplete or inconsistently measured between subjects. An additional advantage of this design is that, since the latency from pre-diabetes to overt diabetes may be longer than the initial 12 months follow-up,23 the long-term follow-up will allow a more accurate estimation of the time trends (cumulative incidences) and clinical features associated with progression to diabetes.

**COLLABORATION**

The Mollerussa study is open to future joint studies with external study groups. Investigators with an interest in hypotheses related to pre-diabetes are welcome to contact a member of the Institut Universitari d’Investigació en Atenció Primària Jordi Gol (IDiAP Jordi Gol) to submit a joint study proposal to the Scientific Committee of the institution. The group will consider these proposals if they are in accordance with the study objectives, and do not overlap with other studies already under way. If accepted, a formal written agreement will be established with the collaborative group.

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**Correction notice** This paper has been amended since it was published Online First. Owing to a scripting error, some of the publisher names in the references were replaced with ‘BMJ Publishing Group’. This only affected the full text version, not the PDF. We have since corrected these errors and the correct publishers have been inserted into the references.

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**Contributors** MF, JRMM and DM conceived and designed the study; MBV, JF-N, and AM participated in the study design; MBV, MF, ER, NA, MG-C, NM and AM collected the data; EC built and managed the database; JRMM managed the database, contributed to data cleaning, performed the statistical analyses and contributed to interpretation of the data; MBV, DM, JF-N and MM-C wrote the manuscript. All authors critically reviewed the manuscript and approved the final version to be published.

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**Competing interests** None declared.

**Patient consent** Obtained.

**Ethics approval** Ethics Committee of the Primary Health Care University Research Institute (IDiAP Jordi Gol) (P12/043).

**Provenance and peer review** Not commissioned; externally peer reviewed.

**Data sharing statement** This article describes the establishment of a longitudinal cohort and early findings, and full results will be submitted for peer-reviewed publication in due course. The authors are willing to share unpublished data with interested parties upon request because they contain identifying human information and are unsuitable for public deposition. Requests may be made to the corresponding author (didacmauricio@gmail.com).

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**REFERENCES**


