

## **CNIC scientists discover a key mechanism in fat cells that protects the body against energetic excess**

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*A study from the CNIC, published in Nature Communications, reveals how caveolae allow adipocytes to expand safely, storing fat without breaking or causing inflammation*

A team at the [Centro Nacional de Investigaciones Cardiovasculares](#) (CNIC), led by Professor [Miguel Ángel del Pozo Barriuso](#), who heads the [Mechanoadaptation and Caveolae Biology](#) group at the CNIC, has identified an essential mechanism in fat cells (adipocytes) that enables them to enlarge safely to store energy. This process avoids tissue damage and protects the body from the toxic effects of accumulating fat molecules (lipids) in inappropriate places. The results, published in '[Nature Communications](#)', signify a major advance in the understanding of metabolic diseases. Moreover, this discovery opens the door to the development of new therapeutic strategies to combat diseases related to chronic energetic excess, such as overweight, obesity, lipodystrophy, and metabolic syndrome, and their grave cardiovascular and metabolic complications.

In modern societies, characterized by sedentary lifestyles and high-calorie diets, adipose tissue is a key determinant of metabolic health. Adipocytes can enlarge to store energy in the form of fat, preventing excess lipids from accumulating in organs like the liver or in the blood vessel wall (especially in the heart and the brain), where they could cause irreparable damage.

Nevertheless, this process is not risk free. When adipocytes are overloaded with fat they can rupture, releasing their toxic contents and generating inflammation and metabolic alterations.

The CNIC study examined how adipocytes adapt to withstand the mechanical stress associated with their expansion to accommodate an excess of fat.

The team analyzed the role of caveolae, small invaginations in the cell membrane that act as sensors and shock absorbers of these stresses. "When an adipocyte accumulates fat and its surface is under increased tensile stress, the caveolae flatten, releasing a 'reservoir' of membrane that allows the cell to enlarge without breaking apart. Conversely, when fat reserves diminish, these structures regroup to reduce the excess membrane and restore cellular stability," explained study first author Dr. María Aboy Parda.

Caveolae: not just a structural support

As well as physically protecting adipocytes, caveolae also play an essential role in coordinating cell metabolism. Miguel Ángel del Pozo Barriuso explained that during adipocyte expansion, "molecular components of these membrane structures travel to other cell compartments, conveying signals that adjust metabolic activity to match the level of energy reserves. This capacity for internal communication makes caveolae key elements for efficient caveolar function."

However, when these structures are absent or malfunction, adipocytes become more rigid, vulnerable to rupture, and less efficient at storing energy. The result, added Aboy Parda, "is an inflammatory reaction that compromises the body's metabolic health. This phenomenon is linked to conditions such as lipodystrophy, in which the body cannot store fat, leading to severe metabolic and cardiovascular alterations."

The CNIC study highlights the key role of the caveolae protein caveolin-1 (Cav-1). For caveolae to flatten correctly in response to fluctuations in mechanical tension in the cell membrane, Cav-1 needs to be chemically altered by the addition of a phosphoryl group to a specific amino acid, a process called phosphorylation. For the study, the researchers developed a transgenic mouse that expresses a genetically altered version of Cav-1 that cannot be phosphorylated. This renders adipocytes unable to expand correctly in response to the mechanical tension generated by the accumulation of lipids (see figure), severely limiting their capacity to store energy and maintain cellular integrity. The failure of this simple mechanism ultimately leads to lipodystrophy and its severe consequences.

Del Pozo Barriuso concluded that "these results give us a better understanding of how adipose tissue

responds to the mechanical forces associated with energetic excess. In the context of obesity and metabolic syndrome, this protective mechanism is essential for minimizing organismal damage.

The study was conducted with the participation of the Transgenesis, Pluripotent Cell Technology, Microscopy, and Histopathology units at the CNIC. The CNIC team also partnered with investigators from the *Instituto de Ciencia de Materiales de Madrid* (ICMM, CSIC), the [Centro de Investigación en Medicina Molecular y Enfermedades Crónicas CIMUS](#) (CIMUS), the *Centro Nacional de Investigaciones Oncológicas* (CNIO), the [Instituto de Investigaciones Biomédicas 'Sols-Morreale'](#), and the [Centro de Investigaciones Biológicas Margarita Salas -CSIC](#).

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