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Submitted to the journal "Open Access Indonesian Journal of Medical Reviews" (March 9th, 2022)

Mechanisms of Cellular Adaptation and Change: A Narrative Literature Review Rachmat Hidayat^{1*}, Catherine²

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Abstract

Injury to cells and the surrounding environment, called the extracellular matrix, triggers injury to tissues and organs. Although a normal cell is limited by narrow structural and functional constraints, it is able to adapt to demands or biological stress to maintain a fixed state called homeostasis. Adaptation is a reversible, structural, or functional response to normal or physiological conditions and adverse or pathological conditions. This review aims to describe the mechanism of cellular adaptation in the human body. Cells adapt to the environment to escape and protect from injury. Cell adaptation be it normal or injury, this condition lies somewhere between these two conditions. The most significant adaptive changes in cells include atrophy (decrease in cell size), hypertrophy (increase in cell size), hyperplasia (increase in cell number), and metaplasia (reversible replacement of one mature cell to another less mature cell or phenotype change). Dysplasia (disorder of cellular growth) is not considered a real cellular adaptation but rather an atypical hyperplasia. In conclusion, cellular adaptation is a central and common part of many disease conditions.

Keywords: Cell adaptation, Cell injury, Molecular, Inflammatory, Metaplasia.

Introduction

Knowledge of the structural and functional reactions of cells and tissues to injury causative agents, including genetic defects, is key to understanding disease processes. Cellular injury can be caused by any factor that disrupts cellular structures or deprives cells of their need for oxygen and nutrients to survive. ¹ Injuries can be reversible (sublethal) or irreversible (lethal) and are broadly classified as chemical, hypoxic (lack of adequate oxygen), free radical, intentional or unintentional, and inflammatory or immunological. ^{2,3} Cellular injuries of varied causes have different clinical and pathophysiological manifestations. Stress from metabolic disorders can be related to intracellular accumulation and includes carbohydrates, proteins, and lipids. Side cell death can lead to calcium accumulation leading to pathological calcification.

Cell death is confirmed by structural changes seen when cells are stained and observed with a microscope. The most important change is the nucleus. Withouta healthy nucleus, the cell cannot survive. The two main types of cell death are necrosis and apoptosis, and nutrient chaos can initiate autophagy leading to cell death. ^{4.5} This review aims to describe the mechanism of cellular adaptation in the human body.

Cell adaptation and injury

Injury to cells and the surrounding environment, called the extracellular matrix, triggers injury to tissues and organs. Although a normal cell is limited by narrow structural and functional constraints, it is able to adapt to demands or biological stress to maintain a fixed state called homeostasis. ¹ Adaptation is a reversible, structural, or functional response to normal or physiological conditions and adverse or pathological conditions. For example, the uterus adapts to pregnancy—a normal physiological condition—with enlargement. Enlargement occurs due to an increase in the size and number of uterine cells. In adverse conditions, such as high blood pressure, myocardial cells are stimulated to enlarge by increasing the work of the pump. Like most of the body's adaptation mechanisms, however, cellular adaptation to adverse conditions is usually only temporarily successful. Bad or long-term stressors overwhelm adaptive processes and cellular injury or death. Changes in cellular and tissue biology can occur from adaptation, injury, neoplasia, accumulation, aging, or death. ^{4,5}

Cellular aging causes structural and functional changes that eventually trigger cell death or a decreased capacity to repair from injury. The mechanisms that explain how and why cells age are unknown, and the difference between the pathological and physiological changes that occur with aging is often elusive. Aging obviously causes changes in cellular structure and function, but aging or getting older is both inevitable and normal.⁶

Adaptationseluler

Cells adapt to the environment to escape and protect from injury. Cell adaptation, be it normal or injury—this condition lies somewhere in between. Adaptation is a reversible change in cell size, number, phenotype, metabolic activity, or cell function. ^{7,8} However, cellular adaptation is a central and common part of many disease conditions. In the early stages of a successful adaptive response, the cell may improve its function; Thus, it is difficult to distinguish pathological responses from extreme adaptation from excessive functional demands. The most significant adaptive changes in cells include atrophy (decrease in cell size), hypertrophy

(increase in cell size), hyperplasia (increase in cell number), and metaplasia (reversible replacement of one mature cell to another less mature cell or phenotype change). ⁹ Dysplasia (disorder of cellular growth) is not considered a real cellular adaptation but rather an atypical hyperplasia. ¹⁰ These changes are shown in figure 1.

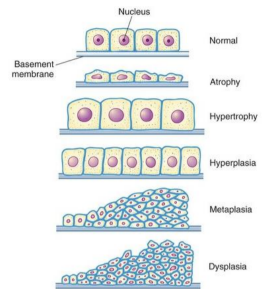


Figure 1. Adaptive changes in simple cuboidal epithelial cells.

Atrophy

Atrophy is a decrease or shrinkage in cellular size. If atrophy occurs in enough organ cells, the entire organ will shrink or become atrophic. Atrophy can affect any organ, but mostly the skeletal muscles, heart, secondary sex organs, and brain (figure 2). Atrophy can be divided into physiological or pathological. Physiological atrophy occurs with early development. For example, the thymus gland undergoes physiological atrophy during childhood. Pathological atrophy occurs as a result of decreased workload, use, pressure, blood flow, nutrition, hormonal stimulation, and nerve stimulation. Individuals who experience immobilization in bed for a long time exhibit skeletal muscle atrophy called disuse atrophy. ¹¹ Aging causes brain cells to become atrophic and endocrine-dependent organs, such as the gonads, to shrink as hormonal stimulation decreases. Whether atrophy is caused by normal physiological conditions or pathological conditions, atrophic cells show the same basis of change.

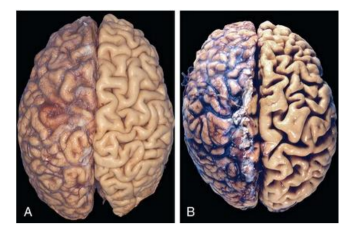


Figure 2. Atrophy. A, The normal brain of young people. B, Brain atrophy in an 82-year-old man with atherosclerosis. Brain atrophy as a result of aging and decreased blood flow. Note that loss of brain substance narrows the gyrus and dilates the sulcus. The meninges have been stripped from the right half of each specimen to reveal the surface of the brain.

Atrophied muscle cells contain less ER and fewer mitochondria and myofilaments (parts of muscle fibers that control contraction) than normal cells do. In muscular atrophy caused by nerve loss, oxygen consumption and amino acid uptake are rapidly reduced. Atrophic mechanisms include decreased potassium synthesis or increased protein degradation, or both. Protein degradation occurs mainly by the ubiquitin-proteosome pathway. ^{10.12}

Atrophy as a result of malnutrition can activate the ubicutin ligase which targets proteins for proteasome degradation. Accelerated protein degradation can be a mechanism responsible for catabolic conditions, including acnker cachexia. Atrophy is often accompanied by a process of "self-eating" called autophagy that induces autophagy vacuoles. These vacuoles are membrane-bound vesicles within the cell that contain cellular debris—small fragments of mitochondria and ER—and hydrolytic enzymes, isolated in autophagy vacuoles to prevent uncontrolled cell destruction. So vacuoles proliferate as necessary to protect uninjured organelles from injured organelles and those eventually taken up and destroyed by lysosomes. The exact content of autophagy vacuoles can resist destruction by lysosomal enzymes and persist in membrane-bound waste bodies. For example, these granules contain lipofuscin, a yellow-brownish aging pigment. Lipofuscin accumulates mainly in liver cells, myocardial cells, and atrophic cells.¹³

Hypertrophy

Hypertrophy is an increase in cell size thereby increasing the size of the affected organ. Much of the knowledge about hypertrophy comes from research on the heart. Cells from the heart

and kidneys are particularly responsive to enlargement. Hypertrophy can be physiological or pathological. Physiological hypertrophy is the result of increased demand, stimulation by hormones (example: atrial natriuretic peptide hormone), and growth factors (example: IGF-1). Physiological hypertrophy in skeletal cells occurs in response to heavy work. Muscular hypertrophy tends to decrease if the excessive workload is also reduced. Pregnancy is an example of physiological hypertrophy and hormone-induced enlargement of the uterus.

Pathological hypertrophy is acquired from chronic hemodynamic overload, for example, from hypertension or dysfunction of the heart valves. A focus of much research is the molecular basis of cardiac hypertrophy because it can develop into maladaptive conditions, including dysrhythmias, heart failure, and sudden death. ¹⁴

Triggers of cardiac hypertrophy include two types of signals: mechanical signals, such as strain, and trophic signals, such as growth factors and vasoactive agents (figure 3). Mechanical strain sensing is triggered from increased workload. This sensor, by itself, can increase the production of growth factors (example: IGF-1) and vasoactive factors (example: angiotensin II). Signals from these membrane sensors activate complex signaling pathways, including phosphoinocytode 3-kinase (PI3K)/AKT pathway and paired G-protein receptors. Transcription factors are activated from signaling pathways to increase muscle protein synthesis. Enlargement of the initials of the heart is caused by dilation of the heart chambers, temporary life, and is followed by increased synthesis of cardiac muscle proteins that enable muscle fibers to work more. The nucleus is also hypertrophied and accentuates increased synthesis of DNA. An increase in cell size is related to an increased accumulation of proteins in cellular components (plasma membrane, ER, myofilaments, mitochondria) and not with an increase in the amount of cell fluid. Over time, cardiac hypertrophy is characterized by remodeling of the extracellular matrix and increased growth of mature myocytes. Prolonged cardiac hypertrophy leads to contractile dysfunction, decompensation, and eventually heart failure. Heart failure is a leading cause of death worldwide. One area of investigation is microRNAs (miRNAs) that regulate the expression of gene targets after transcription. In mice, miRNA 212-/132 regulates cardiac hypertrophy and cardiomyocyte autophagy. Remodeling of heart tissue occurs after cardiac stress and can progress to heart failure and death. Observers are studying the formation of cardiac fibrosis caused by increased activity of cardiac fibroblasts leading to overproduction of the extracellular matrix. Uncoded RNAs (ncRNAs) as gene regulators are one focus for studying cardiac fibrosis and a therapeutic target.¹³

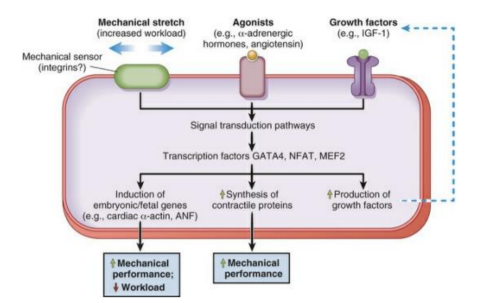


Figure 3. Mechanisms of myocardial hypertrophy. Mechanical sensors seem to be the main stimulators for physiological hypertrophy. Other stimuli that may be more important for pathological hypotrophy include agonists (initiators) and growth factors. This factor then becomes a signal transcription pathway where the transcription factor then binds to the DNA chain, the activation of muscle proteins responsible for hypertrophy. These pathways include embryonal/fetal gene induction, increased contractile protein synthesis, and growth factor production.

Hyperplasia

Hyperplasia is an increase in the number of cells in an organ or tissue resulting from an increase in the cell defense ratio. Hyperplasia occurs in response to an injury that occurs when the wound or injury is aggravating and lasts a long time. ¹⁰ The main mechanism of hyperplasia is the production of growth factors, which stimulate surviving cells (after cell loss or injury) to synthesize new cell components and, eventually, divide. Another mechanism is increased output of new cells from stem cell tissue. For example, if liver cells can be compromised, new cells can regenerate from intrahepatic stem cells. Although hyperplasia and hypertrophy have different processes, they can occur together and the specific mechanism is unknown. Hyperplasia can be physiological or pathological.

Two types of normal, or physiological, hyperplasia are compensatory hyperplasia and hormonal hyperplasia. Compensatory hyperplasia is an adaptive mechanism that enables certain organs to regenerate. For example, removal of the liver part triggers hyperplasia of surviving liver cells (hepatocytes) to compensate for the loss. Even with 70% removal of the liver, complete regeneration can occur in about 2 weeks. Hepar can renew itself by simple duplication of perfectly differentiated cells. Hepatocytes usually live a year or more; Then, through a very slow cell division ratio, they renew themselves. If a large number of hepatocytes are lost from surgery or injury, a burst of cell division occurs from the surviving hepatocytes—

quickly replacing the lost tissue. Much is unknown about stem cell activation and hepatocyte renewal in severe liver injury. ^{5.6}

Significant compensatory hyperplasia occurs in the intestinal epithelium and epidermal, hepatocytes, bone marrow cells, and fibroblasts. An example of compensatory hyperplasia is a callus, or thickening, of the skin as a result of epidermal cell hyperplasia in response to mechanical stimuli. Another example is the response to wound healing as part of the inflammatory process (see Chapter 7).

Hormonal hyperplasia occurs mainly in estrogen-dependent organs, such as the uterus and breasts. After ovulation, for example, estrogen stimulates the endometrium to grow and thickens for reception of the fertilized ovum. If pregnancy occurs, hormonal hyperplasia, such as hypertrophy, enables the uterus to enlarge.

Pathologic hyperplasia is an abnormal proliferation of normal cells and can occur as an excess response to external stimuli or the effects of growth factors on target cells (figure 4). Hyperplastic cells are recognized by enlargement of the nucleus, clumping of chromatin, and the presence of one or more enlarged nucleolus. Most common examples are pathological hyperplasia of the endometrium, which causes an imbalance between estrogen and progesterone with a relative increase in estrogen. Pathologic endometrial hyperplasie, which causes excessive menstrual bleeding, is under the control of regular growth inhibition. If this control fails, endometrial hyperplasia cells can undergo malignant transformation. Benign prostatic hyperplasia is another example of pathological hyperplasia and results from changes in hormonal balance. In both of these instances, if the hormonal imbalance is corrected hyperplasia is reduced. ^{6.7}

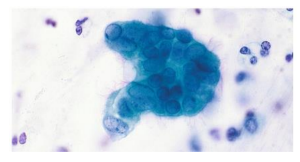


Figure 4. Hiperplasia epithelium bronchi.

Dysplasia

Dysplasia refers to abnormal changes in the size, shape, and arrangement of mature cells. Dysplasia is not considered a real adaptive process but is related to hyperplasia and is often called atypical hyperplasia. Dysplastic changes are mostly found in the epithelium. The architecture of a dysplastic network can be untidy. Importantly, the term dysplasia is not cancer and may not develop into cancer. ⁸ Dysplasia that does not involve the entire thickness of the epithelium may improve completely. Removal of stimulating stimulus, for example, certain hormonal stimuli, in mild to moderate dysplasia that does not involve the entire epithelium may improve. When dysplastic changes penetrate into the basement membrane, it is considered a preinvasive neoplasm and is known as carcinoma in situ.

Metaplasia

Metaplasia is the reversible replacement of one mature cell (epithelial or mesenchymal) with another, sometimes poorly differentiated. Found to be associated with tissue damage, repair, and regeneration. At any given time, the adaptive turnover of cells can better match the changes in their environment. For example, gastroesophageal reflux damages the squamous epithelium of the esophagus, and adaptive changes or alternation by the glandular epithelium may be better tolerated by acidic environments. Usually, however, change is not always beneficial. In long-term smokers, chronic irritation from cigarettes causes ciliated columnar epithelial cells of the trachea and bronchi to be replaced by pseudo-squamous epithelial cells (figure 5). The formed squamous epithelial cells do not secrete mucus or have cilia, causing a loss of vital protective mechanisms. Bronchial metaplasia can be reversible if the induced stimulus, usually smoking, is removed. ⁹ If the induction stimulus remains, it can initiate a malignant transformation of the metaplasia epithelium.

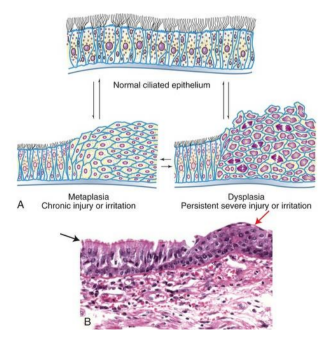


Figure 5. Reversible changes in cell boundaries in the bronchi. A, Normal ciliated epithelium, metaplasia, and dysplasia. B, Histologic view with upper left (black arrow) normal columnar epithelium and basal membrane, and upper right (red arrow) squamous metaplasia.

Metaplasia develops from reprogrammed stem cells and remains in most epithelium or undifferentiated mesenchymal cells (tissue from the embryonic mesoderm) present in connective tissue. ^{8,12} These precursor cells mature along new pathways due to signals generated by cytokines and growth factors in the cell environment. The mechanism of metaplasia is not the result of changes in the phenotype of already differentiated cell types.

Conclusion

Adaptation is a reversible change in cell size, number, phenotype, metabolic activity, or cell function. However, cellular adaptation is a central and common part of many disease conditions.

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Submitted to the journal "Open Access Indonesian Journal of Medical Reviews" (March 9th, 2022)

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Submission acknowledgement

Dear author(s),

Rachmat Hidayat*, Catherine has submitted the manuscript "Mechanisms of Cellular Adaptation and Change: A Narrative Literature Review" to Open Access Indonesian Journal of Medical Reviews. The paper will be screened by editor and reviewed by peer review.

Cordially,



(*) Corresponding author

Peer Review Results "Open Access Indonesian Journal of Medical Reviews (March 9th, 2022)

Open Access Indonesian Journal of Medical Reviews



Peer Review Results

Dear author(s),

Rachmat Hidayat*, Catherine has submitted the manuscript "Mechanisms of Cellular Adaptation and Change: A Narrative Literature Review" to Open Access Indonesian Journal of Medical Reviews. The decision : Revision Required.

Cordially,



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Reviewer 1: Revision required

Mechanisms of Cellular Adaptation and Change: A Narrative Literature Review→1

Rachmat Hidayat^{1*}, Catherine²

¹Department of Biology, Faculty of Medicine, Universitas Sriwijaya, Palembang, Indonesia ²Faculty of Medicine, Universitas Sriwijaya, Palembang, Indonesia *email: rachmathidayat@fk.unsri.ac.id

<mark>Abstract</mark>→3

Injury to cells and the surrounding environment, called the extracellular matrix, triggers injury to tissues and organs. Although a normal cell is limited by narrow structural and functional constraints, it is able to adapt to demands or biological stress to maintain a fixed state called homeostasis. Adaptation is a reversible, structural, or functional response to normal or physiological conditions and adverse or pathological conditions. This review aims to describe the mechanism of cellular adaptation in the human body. Cells adapt to the environment to escape and protect from injury. Cell adaptation be it normal or injury, this condition lies somewhere between these two conditions. The most significant adaptive changes in cells include atrophy (decrease in cell size), hypertrophy (increase in cell size), hyperplasia (increase in cell number), and metaplasia (reversible replacement of one mature cell to another less mature cell or phenotype change). Dysplasia (disorder of cellular growth) is not considered a real cellular adaptation but rather an atypical hyperplasia. In conclusion, cellular adaptation is a central and common part of many disease conditions.

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Introduction→4

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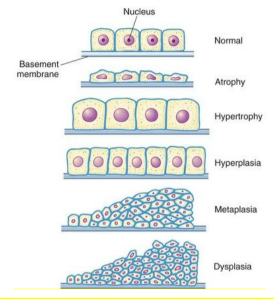


Figure 1. Adaptive changes in simple cuboidal epithelial cells.

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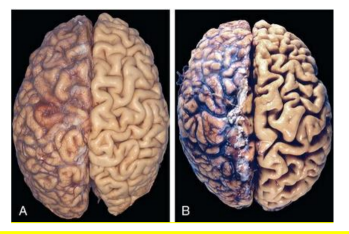


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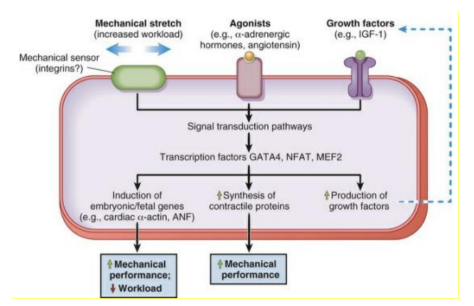


Figure 3. Mechanisms of myocardial hypertrophy. Mechanical sensors seem to be the main stimulators for physiological hypertrophy. Other stimuli that may be more important for pathological hypotrophy include agonists (initiators) and growth factors. This factor then becomes a signal transcription pathway where the transcription factor then binds to the DNA chain, the activation of muscle proteins responsible for hypertrophy. These pathways include embryonal/fetal gene induction, increased contractile protein synthesis, and growth factor production.

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Significant compensatory hyperplasia occurs in the intestinal epithelium and epidermal, hepatocytes, bone marrow cells, and fibroblasts. An example of compensatory hyperplasia is a callus, or thickening, of the skin as a result of epidermal cell hyperplasia in response to mechanical stimuli. Another example is the response to wound healing as part of the inflammatory process (see Chapter 7).

Hormonal hyperplasia occurs mainly in estrogen-dependent organs, such as the uterus and breasts. After ovulation, for example, estrogen stimulates the endometrium to grow and thickens for reception of the fertilized ovum. If pregnancy occurs, hormonal hyperplasia, such as hypertrophy, enables the uterus to enlarge.

Pathologic hyperplasia is an abnormal proliferation of normal cells and can occur as an excess response to external stimuli or the effects of growth factors on target cells (figure 4). Hyperplastic cells are recognized by enlargement of the nucleus, clumping of chromatin, and the presence of one or more enlarged nucleolus. Most common examples are pathological hyperplasia of the endometrium, which causes an imbalance between estrogen and progesterone with a relative increase in estrogen. Pathologic endometrial hyperplasie, which causes excessive menstrual bleeding, is under the control of regular growth inhibition. If this control fails, endometrial hyperplasia cells can undergo malignant transformation. Benign prostatic hyperplasia is another example of pathological hyperplasia and results from changes in hormonal balance. In both of these instances, if the hormonal imbalance is corrected hyperplasia is reduced. ^{6.7}

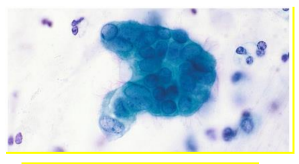


Figure 4. Hiperplasia epithelium bronchi.

<mark>Dysplasia</mark>

Dysplasia refers to abnormal changes in the size, shape, and arrangement of mature cells. Dysplasia is not considered a real adaptive process but is related to hyperplasia and is often called atypical hyperplasia. Dysplastic changes are mostly found in the epithelium. The architecture of a dysplastic network can be untidy. Importantly, the term dysplasia is not cancer and may not develop into cancer. ⁸ Dysplasia that does not involve the entire thickness of the epithelium may improve completely. Removal of stimulating stimulus, for example, certain hormonal stimuli, in mild to moderate dysplasia that does not involve the entire epithelium may improve. When dysplastic changes penetrate into the basement membrane, it is considered a preinvasive neoplasm and is known as carcinoma in situ.

<mark>Metaplasia</mark>

Metaplasia is the reversible replacement of one mature cell (epithelial or mesenchymal) with another, sometimes poorly differentiated. Found to be associated with tissue damage, repair, and regeneration. At any given time, the adaptive turnover of cells can better match the changes in their environment. For example, gastroesophageal reflux damages the squamous epithelium of the esophagus, and adaptive changes or alternation by the glandular epithelium may be better tolerated by acidic environments. Usually, however, change is not always beneficial. In long-term smokers, chronic irritation from cigarettes causes ciliated columnar epithelial cells of the trachea and bronchi to be replaced by pseudo-squamous epithelial cells (figure 5). The formed squamous epithelial cells do not secrete mucus or have cilia, causing a loss of vital protective mechanisms. Bronchial metaplasia can be reversible if the induced stimulus, usually smoking, is removed. ⁹ If the induction stimulus remains, it can initiate a malignant transformation of the metaplasia epithelium.

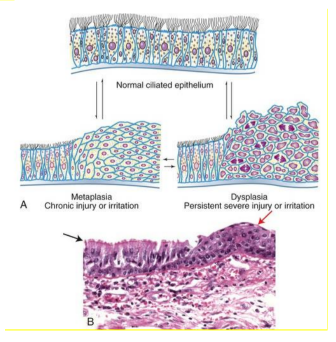


Figure 5. Reversible changes in cell boundaries in the bronchi. A, Normal ciliated epithelium, metaplasia, and dysplasia. B, Histologic view with upper left (black arrow) normal columnar epithelium and basal me mbrane, and upper right (red arrow) squamous metaplasia.

Metaplasia develops from reprogrammed stem cells and remains in most epithelium or undifferentiated mesenchymal cells (tissue from the embryonic mesoderm) present in connective tissue. ^{8,12} These precursor cells mature along new pathways due to signals generated by cytokines and growth factors in the cell environment. The mechanism of metaplasia is not the result of changes in the phenotype of already differentiated cell types.

Conclusion→5

Adaptation is a reversible change in cell size, number, phenotype, metabolic activity, or cell function. However, cellular adaptation is a central and common part of many disease conditions.

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Reviewer Comment:

 $1 \rightarrow$ Title of Manuscripts should be explained main review and declared type of literature review: narrative or systematic review.

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 $5 \rightarrow$ Conclusion should more specific and not more showed more review.

 $6 \rightarrow$ Authors must check the references for make update references. References should no more than 10 years.

Reviewer 2: Revision required

Mechanisms of Cellular Adaptation and Change: A Narrative Literature Review→1 Rachmat Hidayat^{1*}, Catherine²

¹Department of Biology, Faculty of Medicine, Universitas Sriwijaya, Palembang, Indonesia ²Faculty of Medicine, Universitas Sriwijaya, Palembang, Indonesia *email: rachmathidayat@fk.unsri.ac.id

<mark>Abstract</mark>→3

Injury to cells and the surrounding environment, called the extracellular matrix, triggers injury to tissues and organs. Although a normal cell is limited by narrow structural and functional constraints, it is able to adapt to demands or biological stress to maintain a fixed state called homeostasis. Adaptation is a reversible, structural, or functional response to normal or physiological conditions and adverse or pathological conditions. This review aims to describe the mechanism of cellular adaptation in the human body. Cells adapt to the environment to escape and protect from injury. Cell adaptation be it normal or injury, this condition lies somewhere between these two conditions. The most significant adaptive changes in cells include atrophy (decrease in cell size), hypertrophy (increase in cell size), hyperplasia (increase in cell number), and metaplasia (reversible replacement of one mature cell to another less mature cell or phenotype change). Dysplasia (disorder of cellular growth) is not considered a real cellular adaptation but rather an atypical hyperplasia. In conclusion, cellular adaptation is a central and common part of many disease conditions.

Keywords: Cell adaptation, Cell injury, Molecular, Inflammatory, Metaplasia. $\rightarrow 2$

Introduction→4

Knowledge of the structural and functional reactions of cells and tissues to injury causative agents, including genetic defects, is key to understanding disease processes. Cellular injury can be caused by any factor that disrupts cellular structures or deprives cells of their need for oxygen and nutrients to survive. ¹ Injuries can be reversible (sublethal) or irreversible (lethal) and are broadly classified as chemical, hypoxic (lack of adequate oxygen), free radical, intentional or unintentional, and inflammatory or immunological. ^{2,3} Cellular injuries of varied causes have different clinical and pathophysiological manifestations. Stress from metabolic disorders can be related to intracellular accumulation and includes carbohydrates, proteins, and lipids. Side cell death can lead to calcium accumulation leading to pathological calcification.

Cell death is confirmed by structural changes seen when cells are stained and observed with a microscope. The most important change is the nucleus. Withouta healthy nucleus, the cell cannot survive. The two main types of cell death are necrosis and apoptosis, and nutrient chaos can initiate autophagy leading to cell death. ^{4.5} This review aims to describe the mechanism of cellular adaptation in the human body.

Cell adaptation and injury

Injury to cells and the surrounding environment, called the extracellular matrix, triggers injury to tissues and organs. Although a normal cell is limited by narrow structural and functional constraints, it is able to adapt to demands or biological stress to maintain a fixed state called homeostasis. ¹ Adaptation is a reversible, structural, or functional response to normal or physiological conditions and adverse or pathological conditions. For example, the uterus adapts to pregnancy—a normal physiological condition—with enlargement. Enlargement occurs due to an increase in the size and number of uterine cells. In adverse conditions, such as high blood pressure, myocardial cells are stimulated to enlarge by increasing the work of the pump. Like most of the body's adaptation mechanisms, however, cellular adaptation to adverse conditions is usually only temporarily successful. Bad or long-term stressors overwhelm adaptive processes and cellular injury or death. Changes in cellular and tissue biology can occur from adaptation, injury, neoplasia, accumulation, aging, or death. ^{4,5}

Cellular aging causes structural and functional changes that eventually trigger cell death or a decreased capacity to repair from injury. The mechanisms that explain how and why cells age are unknown, and the difference between the pathological and physiological changes that occur with aging is often elusive. Aging obviously causes changes in cellular structure and function, but aging or getting older is both inevitable and normal.⁶

Adaptationseluler

Cells adapt to the environment to escape and protect from injury. Cell adaptation, be it normal or injury—this condition lies somewhere in between. Adaptation is a reversible change in cell size, number, phenotype, metabolic activity, or cell function. ^{7,8} However, cellular adaptation is a central and common part of many disease conditions. In the early stages of a successful adaptive response, the cell may improve its function; Thus, it is difficult to distinguish pathological responses from extreme adaptation from excessive functional demands. The most significant adaptive changes in cells include atrophy (decrease in cell size), hypertrophy

(increase in cell size), hyperplasia (increase in cell number), and metaplasia (reversible replacement of one mature cell to another less mature cell or phenotype change). ⁹ Dysplasia (disorder of cellular growth) is not considered a real cellular adaptation but rather an atypical hyperplasia. ¹⁰ These changes are shown in figure 1.

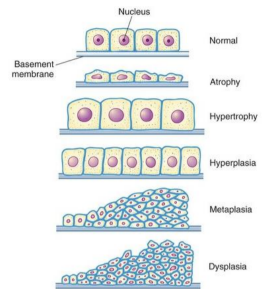


Figure 1. Adaptive changes in simple cuboidal epithelial cells.

Atrophy

Atrophy is a decrease or shrinkage in cellular size. If atrophy occurs in enough organ cells, the entire organ will shrink or become atrophic. Atrophy can affect any organ, but mostly the skeletal muscles, heart, secondary sex organs, and brain (figure 2). Atrophy can be divided into physiological or pathological. Physiological atrophy occurs with early development. For example, the thymus gland undergoes physiological atrophy during childhood. Pathological atrophy occurs as a result of decreased workload, use, pressure, blood flow, nutrition, hormonal stimulation, and nerve stimulation. Individuals who experience immobilization in bed for a long time exhibit skeletal muscle atrophy called disuse atrophy. ¹¹ Aging causes brain cells to become atrophic and endocrine-dependent organs, such as the gonads, to shrink as hormonal stimulation decreases. Whether atrophy is caused by normal physiological conditions or pathological conditions, atrophic cells show the same basis of change.

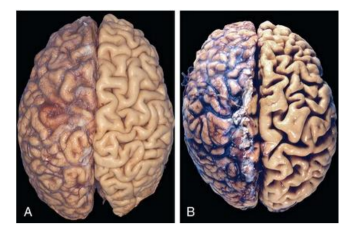


Figure 2. Atrophy. A, The normal brain of young people. B, Brain atrophy in an 82-year-old man with atherosclerosis. Brain atrophy as a result of aging and decreased blood flow. Note that loss of brain substance narrows the gyrus and dilates the sulcus. The meninges have been stripped from the right half of each specimen to reveal the surface of the brain.

Atrophied muscle cells contain less ER and fewer mitochondria and myofilaments (parts of muscle fibers that control contraction) than normal cells do. In muscular atrophy caused by nerve loss, oxygen consumption and amino acid uptake are rapidly reduced. Atrophic mechanisms include decreased potassium synthesis or increased protein degradation, or both. Protein degradation occurs mainly by the ubiquitin-proteosome pathway. ^{10.12}

Atrophy as a result of malnutrition can activate the ubicutin ligase which targets proteins for proteasome degradation. Accelerated protein degradation can be a mechanism responsible for catabolic conditions, including acnker cachexia. Atrophy is often accompanied by a process of "self-eating" called autophagy that induces autophagy vacuoles. These vacuoles are membrane-bound vesicles within the cell that contain cellular debris—small fragments of mitochondria and ER—and hydrolytic enzymes, isolated in autophagy vacuoles to prevent uncontrolled cell destruction. So vacuoles proliferate as necessary to protect uninjured organelles from injured organelles and those eventually taken up and destroyed by lysosomes. The exact content of autophagy vacuoles can resist destruction by lysosomal enzymes and persist in membrane-bound waste bodies. For example, these granules contain lipofuscin, a yellow-brownish aging pigment. Lipofuscin accumulates mainly in liver cells, myocardial cells, and atrophic cells.¹³

Hypertrophy

Hypertrophy is an increase in cell size thereby increasing the size of the affected organ. Much of the knowledge about hypertrophy comes from research on the heart. Cells from the heart

and kidneys are particularly responsive to enlargement. Hypertrophy can be physiological or pathological. Physiological hypertrophy is the result of increased demand, stimulation by hormones (example: atrial natriuretic peptide hormone), and growth factors (example: IGF-1). Physiological hypertrophy in skeletal cells occurs in response to heavy work. Muscular hypertrophy tends to decrease if the excessive workload is also reduced. Pregnancy is an example of physiological hypertrophy and hormone-induced enlargement of the uterus.

Pathological hypertrophy is acquired from chronic hemodynamic overload, for example, from hypertension or dysfunction of the heart valves. A focus of much research is the molecular basis of cardiac hypertrophy because it can develop into maladaptive conditions, including dysrhythmias, heart failure, and sudden death. ¹⁴

Triggers of cardiac hypertrophy include two types of signals: mechanical signals, such as strain, and trophic signals, such as growth factors and vasoactive agents (figure 3). Mechanical strain sensing is triggered from increased workload. This sensor, by itself, can increase the production of growth factors (example: IGF-1) and vasoactive factors (example: angiotensin II). Signals from these membrane sensors activate complex signaling pathways, including phosphoinocytode 3-kinase (PI3K)/AKT pathway and paired G-protein receptors. Transcription factors are activated from signaling pathways to increase muscle protein synthesis. Enlargement of the initials of the heart is caused by dilation of the heart chambers, temporary life, and is followed by increased synthesis of cardiac muscle proteins that enable muscle fibers to work more. The nucleus is also hypertrophied and accentuates increased synthesis of DNA. An increase in cell size is related to an increased accumulation of proteins in cellular components (plasma membrane, ER, myofilaments, mitochondria) and not with an increase in the amount of cell fluid. Over time, cardiac hypertrophy is characterized by remodeling of the extracellular matrix and increased growth of mature myocytes. Prolonged cardiac hypertrophy leads to contractile dysfunction, decompensation, and eventually heart failure. Heart failure is a leading cause of death worldwide. One area of investigation is microRNAs (miRNAs) that regulate the expression of gene targets after transcription. In mice, miRNA 212-/132 regulates cardiac hypertrophy and cardiomyocyte autophagy. Remodeling of heart tissue occurs after cardiac stress and can progress to heart failure and death. Observers are studying the formation of cardiac fibrosis caused by increased activity of cardiac fibroblasts leading to overproduction of the extracellular matrix. Uncoded RNAs (ncRNAs) as gene regulators are one focus for studying cardiac fibrosis and a therapeutic target.¹³

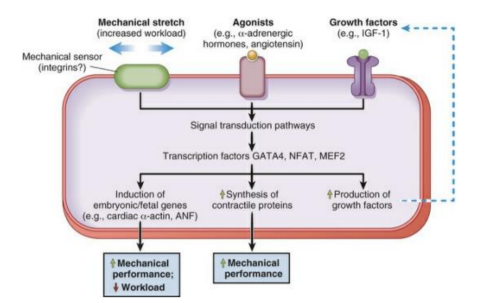


Figure 3. Mechanisms of myocardial hypertrophy. Mechanical sensors seem to be the main stimulators for physiological hypertrophy. Other stimuli that may be more important for pathological hypotrophy include agonists (initiators) and growth factors. This factor then becomes a signal transcription pathway where the transcription factor then binds to the DNA chain, the activation of muscle proteins responsible for hypertrophy. These pathways include embryonal/fetal gene induction, increased contractile protein synthesis, and growth factor production.

Hyperplasia

Hyperplasia is an increase in the number of cells in an organ or tissue resulting from an increase in the cell defense ratio. Hyperplasia occurs in response to an injury that occurs when the wound or injury is aggravating and lasts a long time. ¹⁰ The main mechanism of hyperplasia is the production of growth factors, which stimulate surviving cells (after cell loss or injury) to synthesize new cell components and, eventually, divide. Another mechanism is increased output of new cells from stem cell tissue. For example, if liver cells can be compromised, new cells can regenerate from intrahepatic stem cells. Although hyperplasia and hypertrophy have different processes, they can occur together and the specific mechanism is unknown. Hyperplasia can be physiological or pathological.

Two types of normal, or physiological, hyperplasia are compensatory hyperplasia and hormonal hyperplasia. Compensatory hyperplasia is an adaptive mechanism that enables certain organs to regenerate. For example, removal of the liver part triggers hyperplasia of surviving liver cells (hepatocytes) to compensate for the loss. Even with 70% removal of the liver, complete regeneration can occur in about 2 weeks. Hepar can renew itself by simple duplication of perfectly differentiated cells. Hepatocytes usually live a year or more; Then, through a very slow cell division ratio, they renew themselves. If a large number of hepatocytes are lost from surgery or injury, a burst of cell division occurs from the surviving hepatocytes—

quickly replacing the lost tissue. Much is unknown about stem cell activation and hepatocyte renewal in severe liver injury. ^{5.6}

Significant compensatory hyperplasia occurs in the intestinal epithelium and epidermal, hepatocytes, bone marrow cells, and fibroblasts. An example of compensatory hyperplasia is a callus, or thickening, of the skin as a result of epidermal cell hyperplasia in response to mechanical stimuli. Another example is the response to wound healing as part of the inflammatory process (see Chapter 7).

Hormonal hyperplasia occurs mainly in estrogen-dependent organs, such as the uterus and breasts. After ovulation, for example, estrogen stimulates the endometrium to grow and thickens for reception of the fertilized ovum. If pregnancy occurs, hormonal hyperplasia, such as hypertrophy, enables the uterus to enlarge.

Pathologic hyperplasia is an abnormal proliferation of normal cells and can occur as an excess response to external stimuli or the effects of growth factors on target cells (figure 4). Hyperplastic cells are recognized by enlargement of the nucleus, clumping of chromatin, and the presence of one or more enlarged nucleolus. Most common examples are pathological hyperplasia of the endometrium, which causes an imbalance between estrogen and progesterone with a relative increase in estrogen. Pathologic endometrial hyperplasie, which causes excessive menstrual bleeding, is under the control of regular growth inhibition. If this control fails, endometrial hyperplasia cells can undergo malignant transformation. Benign prostatic hyperplasia is another example of pathological hyperplasia and results from changes in hormonal balance. In both of these instances, if the hormonal imbalance is corrected hyperplasia is reduced. ^{6.7}

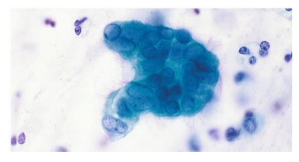


Figure 4. Hiperplasia epithelium bronchi.

Dysplasia

Dysplasia refers to abnormal changes in the size, shape, and arrangement of mature cells. Dysplasia is not considered a real adaptive process but is related to hyperplasia and is often called atypical hyperplasia. Dysplastic changes are mostly found in the epithelium. The architecture of a dysplastic network can be untidy. Importantly, the term dysplasia is not cancer and may not develop into cancer. ⁸ Dysplasia that does not involve the entire thickness of the epithelium may improve completely. Removal of stimulating stimulus, for example, certain hormonal stimuli, in mild to moderate dysplasia that does not involve the entire epithelium may improve. When dysplastic changes penetrate into the basement membrane, it is considered a preinvasive neoplasm and is known as carcinoma in situ.

Metaplasia

Metaplasia is the reversible replacement of one mature cell (epithelial or mesenchymal) with another, sometimes poorly differentiated. Found to be associated with tissue damage, repair, and regeneration. At any given time, the adaptive turnover of cells can better match the changes in their environment. For example, gastroesophageal reflux damages the squamous epithelium of the esophagus, and adaptive changes or alternation by the glandular epithelium may be better tolerated by acidic environments. Usually, however, change is not always beneficial. In long-term smokers, chronic irritation from cigarettes causes ciliated columnar epithelial cells of the trachea and bronchi to be replaced by pseudo-squamous epithelial cells (figure 5). The formed squamous epithelial cells do not secrete mucus or have cilia, causing a loss of vital protective mechanisms. Bronchial metaplasia can be reversible if the induced stimulus, usually smoking, is removed. ⁹ If the induction stimulus remains, it can initiate a malignant transformation of the metaplasia epithelium.

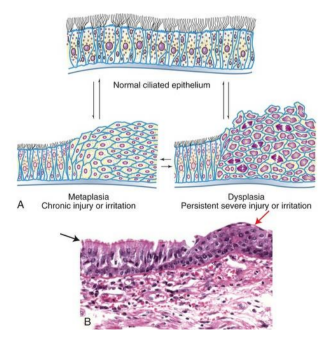


Figure 5. Reversible changes in cell boundaries in the bronchi. A, Normal ciliated epithelium, metaplasia, and dysplasia. B, Histologic view with upper left (black arrow) normal columnar epithelium and basal membrane, and upper right (red arrow) squamous metaplasia.

Metaplasia develops from reprogrammed stem cells and remains in most epithelium or undifferentiated mesenchymal cells (tissue from the embryonic mesoderm) present in connective tissue. ^{8,12} These precursor cells mature along new pathways due to signals generated by cytokines and growth factors in the cell environment. The mechanism of metaplasia is not the result of changes in the phenotype of already differentiated cell types.

Conclusion→5

Adaptation is a reversible change in cell size, number, phenotype, metabolic activity, or cell function. However, cellular adaptation is a central and common part of many disease conditions.

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Mechanisms of Cellular Adaptation and Change: A Narrative Literature Review

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1. Introduction

Knowledge of the structural and functional reactions of cells and tissues to injury-causing agents, including genetic defects, is key to understanding disease processes. Cellular injury can be caused by any factor that disrupts cellular structure or deprives cells of the need for oxygen and nutrients for survival.¹ The injury may be reversible (sublethal) or irreversible (lethal) and is broadly classified as chemical, hypoxic (deficient in adequate oxygen), free radicals, intentional or accidental, and inflammatory or immunological.^{2,3} Cellular injury from a variety of causes own different clinical manifestations and pathophysiology. Stress from metabolic derangements may be related to intracellular accumulation and include carbohydrates, proteins, and lipids. Cell death

ABSTRACT

Injury to cells and the surrounding environment, called the extracellular matrix, triggers injury to tissues and organs. Although a normal cell is limited by narrow boundaries of structure and function, it is capable of adapting to biological demands or stress to maintain a steady state called homeostasis. Adaptation is a reversible, structural, or functional response to normal or physiological conditions and adverse or pathological conditions. This review aimed to describe the mechanism of cellular adaptation in the human body. Cells adapt to the environment to escape and protect against injury. Adaptation of the cell, be it normal or injured, this condition lies somewhere between these two conditions. The most significant adaptive changes in cells include atrophy (decreased cell size), hypertrophy (increased cell size), hyperplasia (increased cell number), and metaplasia (reversible replacement of one mature cell for another less mature cell or change in phenotype). Dysplasia (a disorder of cellular growth) is not considered a true cellular adaptation but rather an atypical hyperplasia. In conclusion, cellular adaptation is a central and common part of many disease conditions.

> sites can lead to calcium accumulation leading to pathological calcification. Cell death was confirmed by visible structural changes when cells were stained and observed under a microscope. The most important change is the nucleus. Without a healthy nucleus, the cell cannot survive. The two main types of cell death are necrosis and apoptosis, and nutrient derangements can initiate autophagy leading to cell death.^{4,5} This review aimed to describe the mechanism of cellular adaptation in the human body.

Cell adaptation and injury

Injury to cells and the surrounding environment, called the extracellular matrix, triggers injury to tissues and organs. Although a normal cell is limited by narrow boundaries of structure and function, it is capable of adapting to biological demands or stress to maintain a steady state called homeostasis.1 Adaptation is a reversible, structural, or functional response to normal or physiological conditions and adverse or pathological conditions. For example, the uterus adapts to pregnancy—a normal physiological condition-by enlargement. Enlargement occurs due to an increase in the size and number of uterine cells. Under adverse conditions, such as high blood pressure, myocardial cells are stimulated to enlarge by increasing their pumping action. Like most of the body's adaptation mechanisms, however, cellular adaptation to adverse conditions is usually only temporary. Adverse or long-term stress overwhelms adaptive processes and causes cellular injury or death. Changes in cellular and tissue biology can occur from adaptation, injury, neoplasia, accumulation, aging, or death.4,5

Cell aging causes structural and functional changes that eventually lead to cell death or decreased capacity to recover from injury. The mechanisms that explain how and why cells age are unknown, and the distinction between the pathological and physiological changes that occur with aging is often elusive. Aging clearly causes changes in cellular structure and function, but aging or growing old is both inevitable and normal.⁶

Cellular adaptation

Cells adapt to the environment to escape and protect against injury. The adaptation of the cell is either normal or injured-the condition lies somewhere in between these two conditions. Adaptation is a reversible change in cell size, number, phenotype, metabolic activity, or cell function.7,8 However, cellular adaptation is a central and common part of many disease conditions. In the early stages of a successful adaptive response, the cell may increase its function; thus, it is difficult to distinguish a pathological response from an adaptation extreme with excessive functional demands. The most significant adaptive changes in cells include atrophy (decreased cell size), hypertrophy (increased cell size), hyperplasia (increased cell number), and metaplasia (reversible replacement of one mature cell for another less mature cell or change in phenotype).9 Dysplasia (derangement of cellular growth) is not considered a true cellular adaptation but rather an atypical hyperplasia.¹⁰ This change is shown in Figure 1.

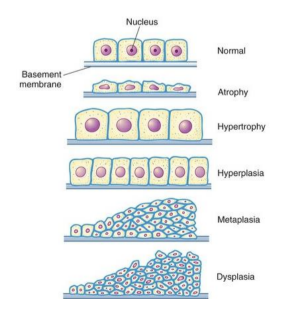


Figure 1. Adaptive changes in simple cuboidal epithelial cells.

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Atrophy is a decrease or shrinkage of cellular size. If atrophy occurs in amount enough organ cells, the entire organ will shrink or become atrophic. Atrophy can affect any organ, but mostly skeletal muscle, heart, secondary sex organs, and brain (Figure 2). Atrophy can be divided into physiological or pathological. Physiological atrophy occurs with early development. For example, the thymus gland undergoes physiological atrophy during childhood. Pathological atrophy occurs as a result of decreased workload, usage, pressure, blood flow, nutrition, hormonal stimulation, and nerve stimulation. Individuals who experience immobilization in bed for a long time show skeletal muscle atrophy which is called disuse atrophy.¹¹ Aging causes brain cells to become surroundings and endocrine-dependent organs, like gonads, to shrink as hormonal stimulation decreases. Whether atrophy is caused by normal physiological conditions or pathological conditions, atrophic cells show the same basic changes.

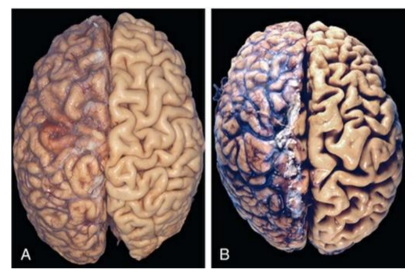


Figure 2. Atrophy. A, The normal brain of a young person. B, Brain atrophy in an 82-year-old man with atherosclerotic disease. Brain atrophy as a result of aging and decreased blood flow. Note that loss of brain substance narrows the gyrus and widens the sulcus. The meninges were stripped from the right half of each specimen to expose the brain surface.

Atrophic muscle cells contain fewer ER and fewer mitochondria and myofilaments (the part of the muscle fiber that controls contraction) than normal cells do. In muscular atrophy caused by nerve loss, oxygen consumption, and amino acid uptake decrease rapidly. The mechanism of atrophy includes decreased synthesis of a protein or increased protein degradation, or both. Protein degradation occurs mainly by the pathway ubiquitin-proteasome.^{10,12}

Atrophy as a result of malnutrition can activate ligase ubiquitin which targets proteins for proteasome degradation. Accelerated protein degradation may be the mechanism responsible for catabolic states, including cachexia cancer. Atrophy is often accompanied by a "self-eating" process called autophagy which induces autophagy vacuoles. These vacuoles are membrane-bound vesicles inside cells that contain cellular debris-small fragments of mitochondria and ER-and hydrolytic enzymes, which are isolated in autophagy vacuoles to prevent uncontrolled cell destruction. Sovacuole proliferates as necessary to protect uninjured organelles from injured organelles and is eventually taken up and destroyed by lysosomes. Certain constituents of autophagic vacuoles can resist destruction by lysosomal enzymes and persist in membrane-bound residual bodies. For example, these granules contain lipofuscin, a yellow-brown aging pigment. Lipofuscin accumulates mainly in hepatic cells, myocardial cells, and atrophic cells.¹³

Hypertrophy

Hypertrophy is an increase in cell size, thereby increasing the size of the affected organ. Much of the knowledge about hypertrophy comes from research on the heart. Cells from the heart and kidneys are particularly responsive to enlargement. Hypertrophy can be physiological or pathological. Physiological hypertrophy is the result caused by increased demand, stimulation by hormones (for example, hormone atrial natriuretic peptide), and growth factors (eg, IGF-1). Physiological hypertrophy of skeletal cells occurs in response to strenuous exercise. Muscular hypertrophy tends to decrease if the excessive workload is also reduced. Pregnancy is an example of physiological hypertrophy and hormone-induced enlargement of the uterus.

Pathological hypertrophy results from chronic hemodynamic overload, for example, from hypertension or valvular dysfunction. A focus of much research is basic molecular from cardiac hypertrophy because it can progress to maladaptive conditions, including dysrhythmias, heart failure, and sudden death.¹⁴

Triggers of cardiac hypertrophy include two types of signals: mechanical signals, such as stretch, and trophic signals, such as growth factors and vasoactive agents (Figure 3). The mechanical strain sensor is triggered by an increase in workload. This sensor, by itself, can increase the production of growth factors (eg, IGF-1) and vasoactive factors (eg angiotensin II). Signals from these membrane sensors activate complex signaling pathways, including the phosphoinositide 3-kinase (PI3K)/AKT pathway and G-protein coupled receptor. Transcription factors are activated from signaling pathways to increase muscle protein synthesis. The initial enlargement of the heart is caused by dilatation of the cardiac chambers, temporary life, and is accompanied by an increase in

cardiac muscle protein synthesis which enables the muscle fibers to work more. The nucleus is also hypertrophied, and features increased synthesis of DNA. The increase in cell size is related to increased protein accumulation in cellular components (plasma membrane, ER, myofilaments, mitochondria) and not to an increase in the amount of cell fluid. Over time cardiac hypertrophy is characterized by extracellular matrix remodeling and increased growth of mature myocytes. Prolonged cardiac hypertrophy pushes contractile dysfunction. decompensation. and eventually heart failure. Heart failure is a leading cause of death worldwide. One area of investigation is microRNA (miRNAs) that regulate target gene expression after transcription. In mice, miRNA 212-/132 regulates cardiac hypertrophy and cardiomyocyte autophagy. Cardiac tissue remodeling occurs after cardiac stress and can progress to heart failure and death. Researchers are currently studying the formation of cardiac fibrosis caused by increased activity of cardiac fibroblasts that lead to the overproduction of extracellular matrix. Non-coding RNAs (ncRNAs) as gene regulators are a focus for studying cardiac fibrosis and therapeutic targets.13

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Two types of normal or physiological hyperplasia are compensatory hyperplasia and hormonal hyperplasia. Compensatory hyperplasia is an adaptive mechanism that enables certain organs to regenerate. For example, the removal of part of the liver triggers hyperplasia of the surviving liver cells (hepatocytes) to compensate for the loss. Even with 70% hepatic removal, complete regeneration can occur in about 2 weeks. The liver is self-renewing by simple duplication of perfectly differentiated cells. Hepatocytes usually live a year or more; then, through a very slow rate of cell division, they renew themselves. If large numbers of hepatocytes are lost from surgery or injury, an explosion of cell division occurs from the surviving hepatocytes—rapidly replacing the lost tissue. Much is not known about stem cell activation and hepatocyte renewal in severe hepatic injury.^{5,6}

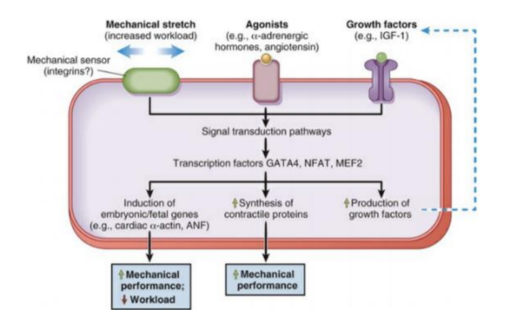


Figure 3. Mechanism of myocardial hypertrophy. Mechanical sensors appear to be the main stimulators for physiological hypertrophy. Other stimuli may be more important to hypertrophy. Pathological factors include agonists (initiators) and growth factors. This factor then becomes a signaling transcription pathway where the transcription factor then binds to the DNA chain, activating muscle proteins responsible for hypertrophy. These pathways include the induction of embryonal/fetal genes, increased contractile protein synthesis, and growth factor production.

Significant compensatory hyperplasia occurs in the intestinal and epidermal epithelium, hepatocytes, bone marrow cells, and fibroblasts. An example of compensatory hyperplasia is a callus, or thickening, of the skin as a result of epidermal cell hyperplasia in response to stimulus mechanics. Another example is the response to wound healing as part of the inflammatory process.

Hormonal hyperplasia occurs mainly in estrogendependent organs, such as the uterus and breasts. After ovulation, for example, estrogen stimulates the endometrium to grow and thicken for the reception of a fertilized ovum. If pregnancy occurs, hormonal hyperplasia, like hypertrophy, enables the uterus to enlarge.

Pathological hyperplasia is an abnormal proliferation of normal cells and can occur in response to excess external stimuli or the effects of growth factors on target cells (Figure 4). Hyperplastic cells are recognized by nuclear enlargement, clumping of chromatin, and the presence of one or more enlarged nucleoli. The most common example is pathological hyperplasia of the endometrium, which causes an imbalance between estrogen and progesterone with a relative increase in estrogen. Hyperplasia pathological endometriosis, which causes excessive menstrual bleeding, is under the control of regular growth restriction. If this control fails, endometrial hyperplastic cells may undergo malignant transformation. Benign prostatic hyperplasia is another example of pathological hyperplasia and results from changes in hormonal balance. In both of these examples, if the hormonal imbalance is corrected, the hyperplasia decreases.^{6,7}

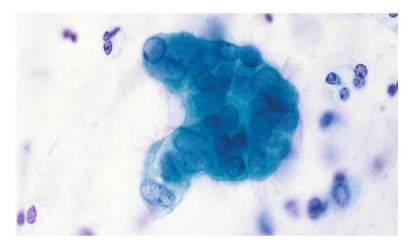


Figure 4. Bronchial epithelial hyperplasia.

Dysplasia

Dysplasia refers to abnormal changes in the size, shape, and arrangement of mature cells. Dysplasia is not considered a true adaptive process but is associated with hyperplasia and is often called atypical hyperplasia. Dysplastic changes are mostly found in the epithelium. The architecture of dysplastic tissue can be untidy. It is also important that the term dysplasia is not cancer and may not develop into cancer.8 Dysplasia that does not involve the full thickness of the epithelium may improve completely. Removal of a stimulating stimulus, for example, certain hormonal stimuli, in mild to moderate dysplasia that does not involve the entire epithelium may be improved. When the dysplastic change penetrates the basement membrane, it is considered a preinvasive neoplasm and is known as carcinoma in situ.

Metaplasia

Metaplasia is the reversible replacement of one mature (epithelial or mesenchymal) cell by another, sometimes less differentiated. Found related to tissue damage, repair, and regeneration. Over time, the adaptive turnover of cells can better match their changing environment. For example, gastroesophageal reflux damages the squamous epithelium of the esophagus, and adaptive changes or replacement by the glandular epithelium may be better tolerated by an acidic environment. Usually, however, change is not always beneficial. In long-term smokers, chronic irritation from smoking causes ciliated columnar epithelial cells of the trachea and bronchi to be replaced by pseudo-squamous epithelial cells (Figure 5). The squamous epithelial cells newly formed do not secrete mucus or have cilia, causing loss of vital protective mechanisms. Bronchial metaplasia may be reversible if the inducing stimulus, usually smoking, is removed.9 If the induction stimulus persists, it can initiate the malignant transformation of the metaplastic epithelium.

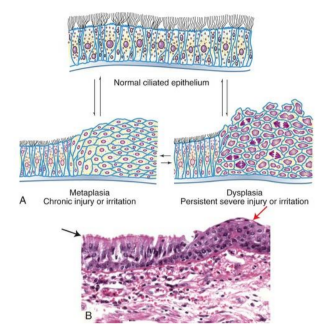


Figure 5. Reversible changes of cell boundaries in bronchi. A, Normal ciliated epithelium, metaplasia, and dysplasia. B, Histological view with top left (black arrow) normal columnar epithelium and basement membrane, and top right (red arrow) squamous metaplasia.

Metaplasia develops from reprogrammed stem cells and persists in most epithelial or mesenchymal cells undifferentiated (tissue from the embryonic mesoderm) present in the connective tissue.^{8,12} These precursor cells mature along new pathways due to signals generated by cytokines and growth factors in the cell environment. The mechanism of metaplasia does not result from a change in the phenotype of a differentiated cell type.

2. Conclusion

Adaptation is a reversible change in cell size, number, phenotype, metabolic activity, or cell function. However, cellular adaptation is a central and common part of many disease conditions.

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Manuscript "Mechanisms of Cellular Adaptation and Change: A Narrative Literature Review" by Rachmat Hidayat*, Catherine, has been accepted to publish in Open Access Indonesian Journal of Medical Reviews Vol 3 issue 2 in April 2023.

Cordially,



(*) Corresponding author

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Mechanisms of Cellular Adaptation and Change: A Narrative Literature Review

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1. Introduction

Knowledge of the structural and functional reactions of cells and tissues to injury-causing agents, including genetic defects, is key to understanding disease processes. Cellular injury can be caused by any factor that disrupts cellular structure or deprives cells of the need for oxygen and nutrients for survival.¹ The injury may be reversible (sublethal) or irreversible (lethal) and is broadly classified as chemical, hypoxic (deficient in adequate oxygen), free radicals, intentional or accidental, and inflammatory or immunological.^{2,3} Cellular injury from a variety of causes own different clinical manifestations and pathophysiology. Stress from metabolic derangements may be related to intracellular accumulation and

ABSTRACT

Injury to cells and the surrounding environment, called the extracellular matrix, triggers injury to tissues and organs. Although a normal cell is limited by narrow boundaries of structure and function, it is capable of adapting to biological demands or stress to maintain a steady state called homeostasis. Adaptation is a reversible, structural, or functional response to normal or physiological conditions and adverse or pathological conditions. This review aimed to describe the mechanism of cellular adaptation in the human body. Cells adapt to the environment to escape and protect against injury. Adaptation of the cell, be it normal or injured, this condition lies somewhere between these two conditions. The most significant adaptive changes in cells include atrophy (decreased cell size), hypertrophy (increased cell size), hyperplasia (increased cell number), and metaplasia (reversible replacement of one mature cell for another less mature cell or change in phenotype). Dysplasia (a disorder of cellular growth) is not considered a true cellular adaptation but rather an atypical hyperplasia. In conclusion, cellular adaptation is a central and common part of many disease conditions.

> include carbohydrates, proteins, and lipids. Cell death sites can lead to calcium accumulation leading to pathological calcification. Cell death was confirmed by visible structural changes when cells were stained and observed under a microscope. The most important change is the nucleus. Without a healthy nucleus, the cell cannot survive. The two main types of cell death are necrosis and apoptosis, and nutrient derangements can initiate autophagy leading to cell death.4,5 This review aimed to describe the mechanism of cellular adaptation in the human body.

Cell adaptation and injury

Injury to cells and the surrounding environment, called the extracellular matrix, triggers injury to

tissues and organs. Although a normal cell is limited by narrow boundaries of structure and function, it is capable of adapting to biological demands or stress to maintain a steady state called homeostasis.¹ Adaptation is a reversible, structural, or functional response to normal or physiological conditions and adverse or pathological conditions. For example, the uterus adapts to pregnancy-a normal physiological condition-by enlargement. Enlargement occurs due to an increase in the size and number of uterine cells. Under adverse conditions, such as high blood pressure, myocardial cells are stimulated to enlarge by increasing their pumping action. Like most of the body's adaptation mechanisms, however, cellular adaptation to adverse conditions is usually only temporary. Adverse or long-term stress overwhelms adaptive processes and causes cellular injury or death. Changes in cellular and tissue biology can occur from adaptation, injury, neoplasia, accumulation, aging, or death.4,5

Cell aging causes structural and functional changes that eventually lead to cell death or decreased capacity to recover from injury. The mechanisms that explain how and why cells age are unknown, and the distinction between the pathological and physiological changes that occur with aging is often elusive. Aging clearly causes changes in cellular structure and function, but aging or growing old is both inevitable and normal.⁶

Cellular adaptation

Cells adapt to the environment to escape and protect against injury. The adaptation of the cell is either normal or injured—the condition lies somewhere in between these two conditions. Adaptation is a reversible change in cell size, number, phenotype, metabolic activity, or cell function.7,8 However, cellular adaptation is a central and common part of many disease conditions. In the early stages of a successful adaptive response, the cell may increase its function; thus, it is difficult to distinguish a pathological response from an adaptation extreme with excessive functional demands. The most significant adaptive changes in cells include atrophy (decreased cell size), hypertrophy (increased cell size), hyperplasia (increased cell number), and metaplasia (reversible replacement of one mature cell for another less mature cell or change in phenotype).⁹ Dysplasia (derangement of cellular growth) is not considered a true cellular adaptation but rather an atypical hyperplasia.¹⁰ This change is shown in Figure 1.

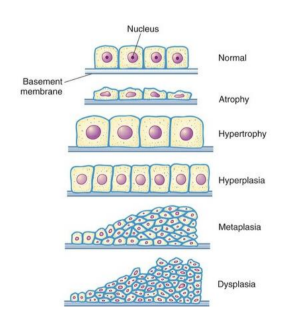


Figure 1. Adaptive changes in simple cuboidal epithelial cells.

Atrophy

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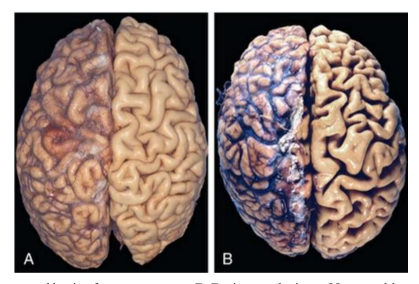


Figure 2. Atrophy. A, The normal brain of a young person. B, Brain atrophy in an 82-year-old man with atherosclerotic disease. Brain atrophy as a result of aging and decreased blood flow. Note that loss of brain substance narrows the gyrus and widens the sulcus. The meninges were stripped from the right half of each specimen to expose the brain surface.

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Triggers of cardiac hypertrophy include two types of signals: mechanical signals, such as stretch, and trophic signals, such as growth factors and vasoactive agents (Figure 3). The mechanical strain sensor is triggered by an increase in workload. This sensor, by itself, can increase the production of growth factors (eg, IGF-1) and vasoactive factors (eg angiotensin II). Signals from these membrane sensors activate pathways, complex signaling including the phosphoinositide 3-kinase (PI3K)/AKT pathway and G-protein coupled receptor. Transcription factors are activated from signaling pathways to increase muscle protein synthesis. The initial enlargement of the heart

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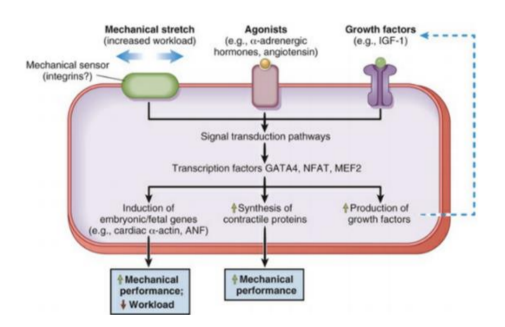


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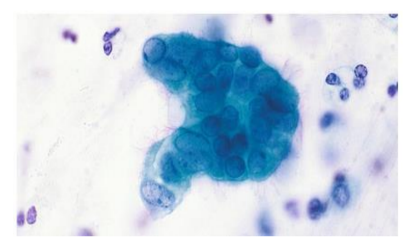


Figure 4. Bronchial epithelial hyperplasia.

Dysplasia

Dysplasia refers to abnormal changes in the size, shape, and arrangement of mature cells. Dysplasia is not considered a true adaptive process but is associated with hyperplasia and is often called atypical hyperplasia. Dysplastic changes are mostly found in the epithelium. The architecture of dysplastic tissue can be untidy. It is also important that the term dysplasia is not cancer and may not develop into cancer.8 Dysplasia that does not involve the full thickness of the epithelium may improve completely. Removal of a stimulating stimulus, for example, certain hormonal stimuli, in mild to moderate dysplasia that does not involve the entire epithelium may be improved. When the dysplastic change penetrates the basement membrane, it is considered a preinvasive neoplasm and is known as carcinoma in situ.

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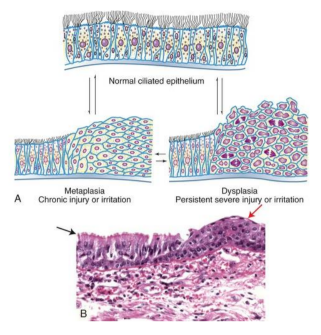


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CERTIFICATE

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