

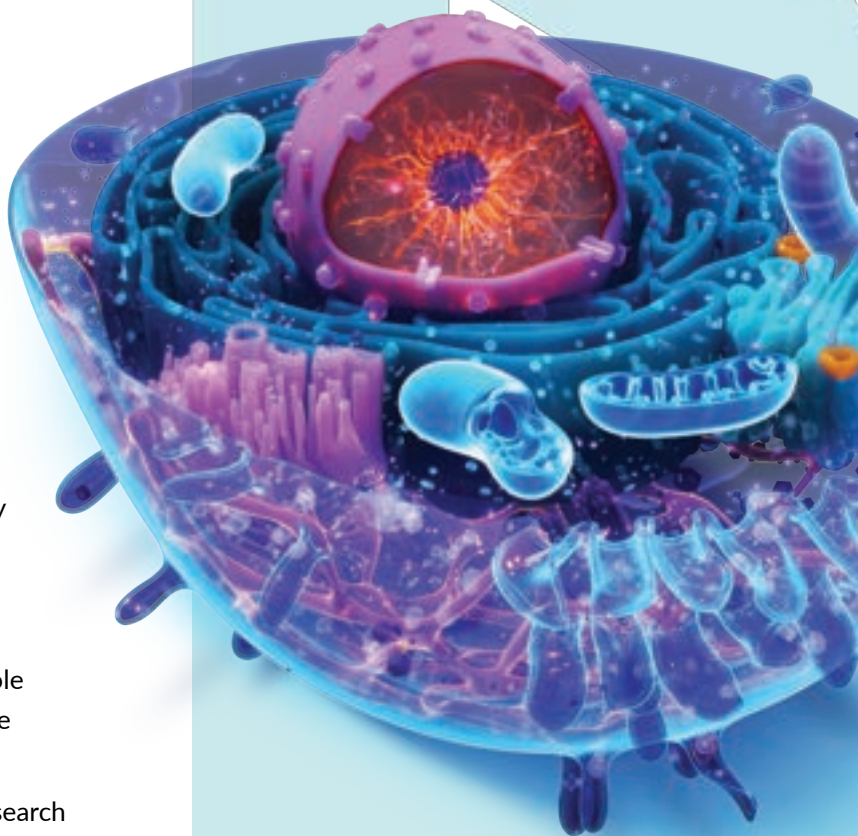
WHITE PAPER

Advancing Mitochondrial Science: Perspectives on Cellular Health and the Role of Ubiquinol

Mitochondria are increasingly understood as central to systemic wellness and healthy aging, helping to coordinate processes such as immune balance, inflammatory control, and cellular repair.^{1,2} Ubiquinol—the antioxidant form of CoQ10—works within the mitochondrial membrane to support these functions through its dual role in mitochondrial energy production and antioxidant defense.^{3,4}

Oxidative stress increases with age, while antioxidant capacity naturally declines. To expand on current knowledge, researchers continue to investigate the role of ubiquinol in maintaining mitochondrial performance and redox balance.^{4,5}

This white paper explores established and emerging research on ubiquinol's role in mitochondrial function. It offers an evidence-based perspective for product developers, formulators, and healthcare professionals evaluating ubiquinol's positioning in systemic wellness applications.



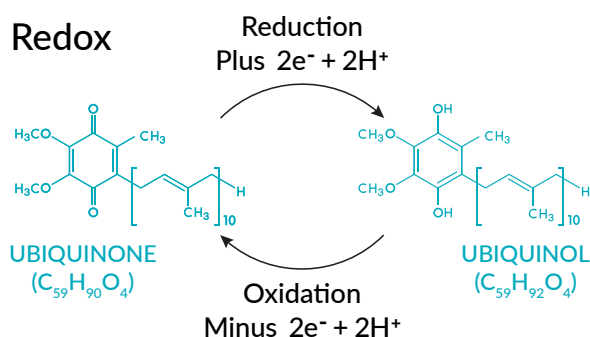
Mitochondria and the Ubiquinol Redox Cycle: Foundations of Cellular Function

Cellular health has long been an area of interest across the health and nutrition industry, since it is recognized as a driver of long-term physiological resilience and healthy aging. New scientific research has brought renewed attention to the mechanisms that influence how cells function and maintain integrity.

One area of expanding scientific inquiry is the mitochondria. While recognized for their role in producing adenosine triphosphate (ATP), the molecular source of energy of the cell, mitochondria are now understood to participate in broader physiological processes, such as coordinating metabolic adaptation, redox signaling, and immune and cellular stress responses, making them integral to maintaining cellular and systemic homeostasis.^{1,5,6}

In tissues with high energy demands, mitochondria are especially dense and functionally active. Each mitochondrion houses thousands of electron transport chains (ETCs), the systems that generate ATP. In this process, electrons are passed along a series of complexes embedded in the mitochondrial inner membrane, creating a proton gradient that drives ATP synthesis.⁷

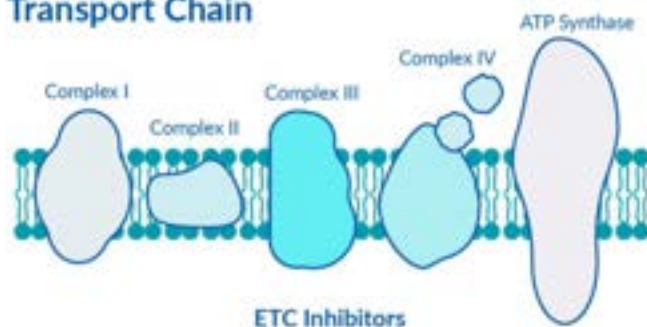
Ubiquinone and ubiquinol participate in the ETC as electron carriers. These two forms of coenzyme Q10 (CoQ10) comprise a redox pair: ubiquinone, its oxidized form, and ubiquinol, the reduced antioxidant form. Their ability to cycle between oxidized and reduced states enables them to transfer electrons between complexes on the ETC.^{3,7}



“Ubiquinone and ubiquinol are flipping back and forth constantly—thousands of times a second—like a light switch on overdrive. That rapid cycling is what keeps the mitochondria humming with energy while keeping free radicals in check.”

— Risa Schulman, Ph.D., President, Tap-Root, Sr. Scientific and Regulatory Advisor to Kaneka Nutrients

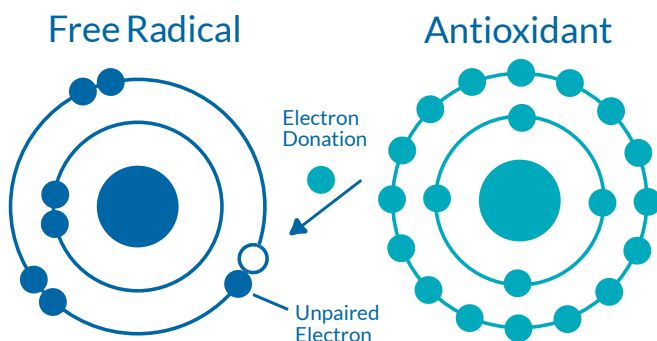
Electron Transport Chain



Some electrons leak from the ETC and react with oxygen, forming reactive oxygen species (ROS), including superoxide and the hydroxyl radical. Under normal conditions, antioxidants neutralize free radical ROS. However, when ROS production exceeds the cell's capacity to neutralize them, oxidative stress can occur, causing damage to cells and tissues.⁴

Mitochondrial membranes and DNA are particularly sensitive to oxidative stress due to their proximity to the ETC. As a result, ETC components, lipid membranes, proteins, and DNA can be affected by excess ROS. This, in turn, can influence ATP synthesis, reduce energy availability, and increase ROS generation—a process that becomes increasingly relevant with age, as antioxidant capacity and mitochondrial efficiency decline.^{8,9}

Ubiquinol's antioxidant activity allows it to play a role in cellular defense by neutralizing ROS. Ubiquinol is lipid soluble and located within the bilayer of the mitochondrial inner membrane—placing it at the source of ROS created by the ETC. Ubiquinol readily donates electrons to neutralize ROS, helping to protect mitochondrial membrane phospholipids and proteins from oxidative modification.⁴



Ubiquinol Is a Unique Antioxidant

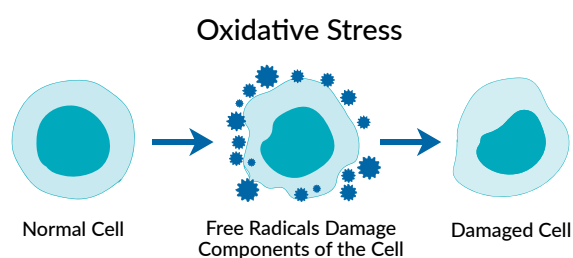
It is the only **lipid-soluble antioxidant** that is endogenously synthesized, **capable of donating two electrons, and continuously regenerated** through enzymatic redox cycling.

It is also **found in almost every cell in the body**, throughout the cells and mitochondrial structures.

These properties make it a **key contributor** to redox homeostasis and proper mitochondrial function.^{3,4}

The Significance of Redox Balance

Redox balance is the steady state between reduction (electron gain) and oxidation (electron loss) reactions within cells. This balance is shaped by both pro-oxidant activity—such as the generation of ROS within the ETC—and antioxidant defense systems that keep those processes in check. When equilibrium is preserved, mitochondria can efficiently produce energy and support essential cellular signaling. When the balance shifts toward oxidation without sufficient antioxidants to manage it, cells experience oxidative stress.^{4,7}



The ratio of ubiquinol to ubiquinone, commonly referred to as the CoQ10 balance, is a measure of oxidative stress. In healthy adults, approximately 95% of the circulating CoQ10 in the blood is in the ubiquinol form.¹⁰ A high percentage of ubiquinol within total CoQ10 reflects a healthy balance; lower blood ubiquinol levels are correlated with increased oxidative stress, partly due to higher utilization.^{11,12}

With age, the body's ability to convert ubiquinone to ubiquinol becomes less efficient, and total circulating CoQ10 shifts toward a more oxidized state.¹³ When the percentage of ubiquinol in the blood decreases, it is an indication that the body is using it as an antioxidant faster than it can be synthesized and replenished. This may affect the mitochondrial antioxidant defenses, increasing susceptibility to oxidative stress and affecting mitochondrial structures and function.^{4,14}

When ubiquinol levels fall, cells may be less able to sustain sufficient energy generation and antioxidant protection. Clinical research shows supplementation can elevate circulating CoQ10 levels and improve the ubiquinol:ubiquinone ratio, supporting healthier redox balance and reducing oxidative stress.^{15,16} Statin use offers a clear example of this relationship, as statins have been shown to lower circulating CoQ10.^{17,18} Research has demonstrated that replenishment through Ubiquinol supplementation increases total CoQ10 levels and is associated with improved muscle discomfort.¹⁹

The Regulatory Reach of Mitochondria

Beyond energy production, mitochondria help regulate cellular processes by responding to changes in redox balance, metabolic demand, and energy status. In doing so, they influence processes such as gene expression, immune signaling, and stress adaptation—all of which may change with age or shifts in cellular demands.^{1,6,20}

One of the distinctive features of mitochondria is their genetic structure. Each contains its own mitochondrial DNA (mtDNA), which encodes a small number of proteins essential for the ETC. Most mitochondrial proteins are encoded by the cell's nuclear DNA, creating a division of genetic responsibility between genomes. Because of the split, communication is required between the nuclear and mitochondrial genomes—sometimes referred to as mitochondrial-nuclear crosstalk—to meet the changing demands of the cell.²¹

“Research has demonstrated that the health of mitochondria and their DNA is critical for the health of cells, tissues, and the body throughout the lifespan. Mitochondrial DNA is located close to where ROS are created, and the structure of mtDNA is tightly packed with a lot of genetic material. This means there is a higher chance of ROS impacting mtDNA. With the important role that mitochondria and their DNA play as ‘CEO of the cell’ (chief executive organelle), keeping ROS in check is so important to maintain the health of mtDNA and cellular homeostasis.”

— Melissa Olivadoti, Ph.D., CMPP, President, Assisi Consulting, Medical Affairs Consultant to Kaneka Nutrients

Mitochondria also serve as sensors. They detect changes in energy substrates, levels of oxidative stress, and shifts in the ATP:ADP ratio—the balance between ATP and its precursor, adenosine diphosphate (ADP). In response to these shifts, mitochondria may adjust energy production, activate antioxidant defenses, trigger signaling pathways that affect nuclear gene expression and protein synthesis, and initiate quality control processes.^{5,22,23}

Ubiquinol helps modulate mitochondrial responses to changing cellular conditions. As oxidative stress increases, due to metabolic activity, aging, or external stressors such as environmental or lifestyle factors, ubiquinol's lipid solubility and location in the inner mitochondrial membrane enable it to efficiently neutralize ROS and protect cells, proteins, and DNA from oxidation.

Ubiquinol's location and role within mitochondria, including its antioxidant capacity, are known to influence mitochondrial function by helping limit ROS accumulation and protect mitochondrial membranes. While these functions and regulatory processes continue to be studied, mitochondria are now known as crucial contributors to cellular regulation and adaptation.⁶

As research continues to uncover the regulatory influence of mitochondria, attention has also turned to what happens when these systems falter. Disruptions in mitochondrial signaling, redox balance, or quality control have been associated with cumulative cellular effects, including impaired ATP production, excess ROS generation, and inappropriate activation of apoptotic pathways. Over time, these changes contribute to the biological processes of aging and may influence how cells respond to metabolic and oxidative challenges.^{23,26}

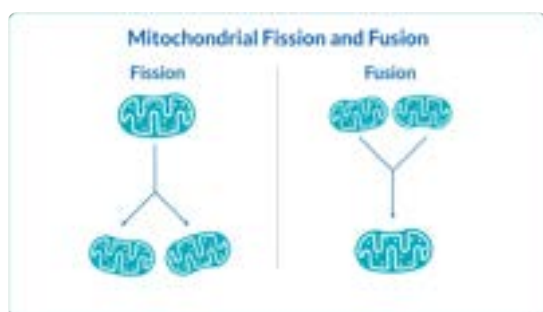
Mitochondrial Quality Control: Fission, Fusion, Mitophagy, Apoptosis, and the Redox Connection

Due to their role in energy production and signaling, mitochondria are constantly producing reactive oxygen species, and if these build up, cells can be subjected to oxidative stress. To maintain cellular health, cells rely on a network of mitochondrial quality control mechanisms that monitor, repair, and, when necessary, remove damaged organelles and cells.^{24,25}

"We're born with all the mitochondria we'll ever have. We can't create new ones from scratch, so the cell has to fix them or blend existing ones together."

— Risa Schulman, Ph.D.

Key components of the mitochondrial quality control system include fission and fusion, mitophagy, and apoptosis. Each of these processes is tightly regulated and responsive to changes in energy status and redox balance.



Fission and fusion are processes that work to preserve mitochondrial function and adapt to cellular needs. Fission allows mitochondria to divide, isolating damaged sections and enabling more efficient distribution of energy production units, improving

cellular energy generation. Fusion merges mitochondrial membranes, allowing the exchange of contents, which helps maintain energy production under increased demand or during repair. When the balance between fission and fusion is disrupted, mitochondrial function is diminished, reducing ATP output and increasing vulnerability to oxidative stress, particularly in tissues with high energy demand, such as the heart and skeletal muscle.^{22,26}

"Fission and fusion are a constant dynamic process of joining and splitting the mitochondria used to repair damage, recycle parts, and keep energy flowing under stress."

— Risa Schulman, Ph.D.

Mitophagy serves as the cellular cleanup process that selectively removes mitochondria that are no longer functioning properly. The best-known pathway uses the proteins PINK1 and Parkin to mark damaged mitochondria for clearance via the cell's recycling system, thereby helping maintain mitochondrial integrity and cellular homeostasis.^{25,27}

Apoptosis, also known as programmed cell death, may be triggered when cellular stress surpasses the capacity for adaptation or repair. A key step in this process is a change in mitochondrial outer membrane permeability, which allows the release of apoptogenic factors—such as cytochrome C and other proapoptotic proteins—into the cytosol, the cell's internal fluid. These signals activate a cascade of enzymes (caspases) that dismantle the cell in a controlled manner.

Apoptosis is generally noninflammatory, allowing damaged cells to be removed without disrupting surrounding tissue.^{24,28}

Redox status plays a central role in regulating these responses. Mitochondrial ROS serve as both stress signals and regulatory cues. At moderate levels, they may promote repair and adaptation; at high levels, they can cause damage to mitochondrial and cellular components, initiate apoptosis, and influence other stress-response pathways.^{1,29}

Research suggests that ubiquinone and ubiquinol may influence aspects of mitochondrial quality control systems, particularly through their redox activity. In in vitro models, exposure to ubiquinone and ubiquinol is associated with:

- Reduced mitochondrial fission and maintenance of mitochondrial structure in oxidative stress conditions³⁰
- Support for the expression of fusion-related proteins, which may enhance mitochondrial adaptability³⁰
- Modulation of mitophagy-related pathways and gene expression, particularly in models with impaired mitochondrial turnover^{31,32}

These effects have been linked to ubiquinol's ability to neutralize ROS. In fact, even when ubiquinone was supplied in vitro, investigators have reported increases in observed antioxidant activity, indicating that ubiquinone converted to ubiquinol. This, in turn, supports the cell's natural adaptive and maintenance processes.

“ROS is the Goldilocks of cell health. If there isn't enough ROS, important healing functions and signaling mechanisms may not take place—for example, muscle building after exercise. With too much, there is damage to the organelles, cells, or tissues. Maintaining a healthy antioxidant level helps to balance ROS levels, including the increase in ROS that also naturally occurs with age.”

— Melissa Olivadoti, Ph.D., CMPP

Mitochondrial Dynamics and Cell Senescence

Mitochondria manage cellular communication pathways that influence immune signaling, cell identity, and cell fate. One way mitochondria contribute to immune activity is through the controlled generation of ROS, which act as secondary messengers in immune pathways that can activate transcription factors such as NF-κB and promote assembly of innate immune complexes. While these processes are essential to cellular defense, prolonged redox imbalance can disrupt immune signaling.²

Senescence is a nonproliferative state that cells may enter in response to significant stress or damage. Senescent cells remain metabolically active and release a set of signaling molecules referred to as the senescence-associated secretory phenotype (SASP), which can affect nearby cells and tissues. While temporary senescence is considered part of a normal stress response, reduced clearance of senescent cells over time has been associated with age-related shifts in tissue function.^{20,33}

“Senescence is like a cell being put on probation. The cell essentially gets sidelined, and if it sits there too long, it can start making trouble—secreting inflammatory signals that affect nearby tissues and even the whole body. Mitochondria are part of the oversight system, helping regulate whether those cells stay in check or spiral out of control.”

— Risa Schulman, Ph.D.

In vitro studies using cultured human endothelial and dermal fibroblast cells have reported reduced ROS levels and positive changes in SASP expression following ubiquinol exposure.³⁴⁻³⁶

Formulary Applications of Ubiquinol: Areas of Interest and Scientific Considerations

As scientific understanding of mitochondrial function expands, ubiquinol is drawing increased interest from product developers, formulators, and healthcare professionals seeking to support cellular wellness and mitochondrial efficiency.

Research demonstrates that ubiquinol is a vital nutrient that supports health and wellness across adult life stages, underscoring the importance of maintaining healthy ubiquinol levels to promote cardiovascular health, healthy aging, preconception health, and menopausal wellness, and support exercise and physical performance.

Bioavailability

Kaneka Ubiquinol® is two times better absorbed than a conventional CoQ10 (ubiquinone) supplement³⁷ and does not require conversion to function as an antioxidant.^{15,38} Research demonstrates that 200 mg of Kaneka Ubiquinol® increases ubiquinol levels by approximately eight times compared to baseline in healthy adults when taken daily for at least 30 days.¹⁶

“Despite ubiquinol being a form of CoQ10, it is molecularly unique compared to the ubiquinone form of CoQ10, and the two forms have different jobs in the body. This is especially important as the body ages, because it becomes harder to make ubiquinol from CoQ10 (ubiquinone), but ubiquinol naturally reverts to CoQ10 after it does its job as an antioxidant. If it's the antioxidant form you want in a supplement formulation, Ubiquinol should be your form of choice.”

— Melissa Olivadoti, Ph.D., CMPP

Cardiovascular Support

Ubiquinol promotes heart health by supporting the high energy requirements of the heart⁷ and protecting cells from damage caused by oxidative stress.³⁹

Clinical research shows that Ubiquinol supplementation improves certain blood markers associated with heart health and that ubiquinol status is associated with heart health.^{40,41} Additional studies highlight the relevance of these markers as indicators associated with cardiovascular health.^{42,43}

Ubiquinol has been shown in a clinical study to benefit vessel health in the following ways¹⁵:

- Facilitating proper vasodilation and blood circulation
- Enhancing nitric oxide production
- Protecting low-density lipoprotein (LDL) from oxidation, which promotes vascular health

In addition, ubiquinol helps to maintain mitochondrial membrane integrity, which is now understood to be foundational for maintaining heart and blood vessel structure and function, particularly with age.⁴⁴⁻⁴⁶

“When we think heart health, we often think of heart rate and blood pressure. But heart health includes the wear and tear on blood vessels and how nutrients reach tissues in the farthest parts of the body, including the brain. If tissues are to remain healthy as the body ages, heart health is a key part of the equation.”

— Melissa Olivadoti, Ph.D., CMPP

Healthy Aging

Clinical studies show that supplementation with Kaneka Ubiquinol[®] increases plasma ubiquinol levels.¹⁵ Also, research demonstrates a positive association between total CoQ10 status, percentage of plasma ubiquinol, and overall physical function in older adults.^{47,48} Higher blood ubiquinol levels also promote cardiovascular health¹⁵ and muscle health.⁴⁷

Preconception Health

Healthy mitochondrial function is essential for the proper function of reproductive cells,⁴⁹ which have high energy demands⁵⁰ and are sensitive to oxidative stress. Age-related changes in oxidative stress and mitochondrial function have been associated with changes in sperm health,⁵¹ as well as oocyte function and ovarian reserve—particularly in women over 35.⁵²

Female preconception support: Ubiquinol supports mitochondrial function and cellular energy generation—essential for oocyte quality and overall egg health.⁵²⁻⁵⁴

Male preconception support: Kaneka Ubiquinol[®] supports healthy sperm function, including morphology, concentration, and motility,^{55,56} and provides antioxidant protection to seminal fluid.⁵⁷

Menopausal Wellness

During perimenopause and menopause, the gradual decline of estrogen, which is an antioxidant, contributes to the common physical, social, and psychological changes that women experience during this time.⁵⁸ Research demonstrates that postmenopausal women exhibit higher oxidative stress than premenopausal women.⁵⁹⁻⁶¹

Kaneka Ubiquinol® supports general health and well-being during and after menopause. In a consumer use study, 80% of menopausal women taking 200 mg of Kaneka Ubiquinol® per day reported decreased irritability, sensitivity, stress, and mood swings after 60 days of supplementation.^{62,63}

“As ubiquinol levels decline with age, losing another powerful antioxidant in estrogen during menopause can be a double-whammy to the body. It’s not surprising that ROS increase during this period, and conditions of aging can feel like they hit warp speed after this change. Ubiquinol is a great option for supplementing antioxidant balance during this time.”

— Melissa Olivadoti, Ph.D., CMPP

Exercise and Physical Performance

During exercise, contraction of the heart and muscles increases, upping the demand for ATP as much as 10-fold.⁶⁴ This is why heart muscle cells contain three to five times more mitochondria than most other cells⁶⁴ and other muscle cells are dense with mitochondria in specialized arrangements for maximal ATP generation and access. The increase in ATP generation also elevates the production of ROS, increasing oxidative stress during and after exercise.⁶⁵

Additionally, athletic ability is influenced by training intensity, age, and baseline fitness,⁶⁶ as well as stressors such as high altitude. Exercising at high

altitude can further challenge oxidative balance by increasing hypoxia-driven ROS production and placing an additional strain on oxygen delivery and cardiorespiratory capacity.⁶⁷

Research shows that Kaneka Ubiquinol® supports exercise and physical performance by:

- Promoting a healthy oxidative balance during exercise⁶⁸
- Supporting energy metabolism during exercise⁶⁹
- Helping the body adapt to rigorous exercise^{70,71}
- Enhancing peak power production in elite athletes when taken at 300 mg per day⁷²
- Promoting healthy cardiac function at high altitude^{70,71}

In a clinical study, participants taking 200 mg of Kaneka Ubiquinol® per day reduced the overall decrease in cardiorespiratory fitness experienced when exercising at high altitude by almost half compared to the placebo—a decrease of 11% in the Ubiquinol group versus 21% for the placebo group.⁷⁰

“Anyone who has tried to sprint through the Denver airport to grab a tight connection can attest to the effects of altitude on exercise. Ubiquinol is a great way to manage the increase in ROS and demands of altitude on the cardiovascular and respiratory system.”

— Melissa Olivadoti, Ph.D., CMPP

High-Altitude Acclimatization: Clinical Observation

In a randomized, placebo-controlled clinical trial conducted during high-altitude acclimatization, participants receiving 200 mg of Ubiquinol daily for approximately six weeks reported reduced fatigue compared with placebo.⁷¹

The Kaneka Nutrients Advantage

Kaneka Nutrients is the manufacturer of Kaneka Ubiquinol®, the leading Ubiquinol supplement, offering unmatched quality, bioavailability, and scientific and clinical data. A partnership with the Kaneka Nutrients team provides the following advantages:

- **Dedication to scientific research:** For over 50 years, Kaneka has invested in researching ubiquinone and ubiquinol, demonstrating our commitment to scientific validation. Kaneka Ubiquinol® is backed by rigorous scientific research and clinical experience across numerous health areas, confirming its safety, efficacy, and health benefits. Currently, there are more than 100 clinical studies on Kaneka Ubiquinol® and over 18 years of positive consumer experience with Kaneka Ubiquinol® supplementation.
- **Patented manufacturing process:** Kaneka Ubiquinol® is a proprietary form of CoQ10 made by utilizing non-genetically modified natural yeast and a patented microbiological fermentation process. This method ensures the production of the all-trans isomer of Ubiquinol, which is bioidentical to the ubiquinol naturally produced in the human body.
- Every stage of production follows FDA regulations and current Good Manufacturing Practices (cGMP), with strict quality assurance processes. This commitment to manufacturing excellence, combined with our deep scientific expertise, decades of research, proprietary data on the CoQ10 molecule, and extensive testing, means Kaneka Ubiquinol® is unparalleled in the marketplace. Our approach yields a pure, high-quality product—free from the cis-isomer generated from synthetic CoQ10 production.

- **Made in the USA:** As the trusted leader in Ubiquinol production in the United States, Kaneka Nutrients ensures the quality, reliability, and purity of our Kaneka Ubiquinol®, delivering the safety and performance you need to enhance your premium product formulations.

**Partner with
Kaneka Nutrients, the
leader in Ubiquinol
innovation, to support
cellular health and
deliver solutions that
promote wellness.**

Get the Kaneka Ubiquinol® Advantage

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