The Breathing-Book
Spontaneous breathing during artificial ventilation

Ernst Bahns
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Introduction

Modern ventilation systems offer various measures for the treatment of respiratory disorders. Today these devices do not only enable sufficient respiratory gas delivery for keeping up the gas exchange in the lung; thanks to advanced pneumatics, electronics and especially due to the implementation of computer technology, artificial ventilation may nowadays be specifically adapted to the respective gas exchange disorder.

Artificial ventilation still remains to be a serious intervention for the patient. Specifically, the pressure conditions generated in the process differ considerably from those during natural breathing. These mechanical pressures affect the lungs and cardiovascular system which may result in undesired side effects. The goal to keep up the vital function of gas exchange with artificial breathing may thus be achieved only by reaching a balance of risks and benefits.

Modern ventilation still faces several partially contradictory challenges. It must serve as a life preserving function with modes that causes stress on the patient however at same time it must keep side effects to a minimum. Modern ventilation faces the contradictory challenges with a new self-conception: The artificial measure is no longer the focus, but rather the natural function of normal breathing even though malfunctioning.
The ventilation system must adapt to the patient and not vice versa. Disturbance of spontaneous breathing by artificial ventilation must be reduced to the absolute minimum. Spontaneous breathing must be promoted whenever possible. This applies not only to the period of weaning, but also to the complete process of artificial ventilation.

We have known about ventilation procedures that allow spontaneous breathing at any time since the 1980’s. We introduced the first mode of this kind in our book „BIPAP/PCV+ - two steps forward in intensive ventilation“. Meanwhile, intensive ventilation has progressed by a few additional steps. The goal of this introduction is to describe the progress and thus highlight the relevance of spontaneous breathing during artificial ventilation.

Hence the title: „The Breathing-Book“
The significance of spontaneous breathing

SPONTANEOUS BREATHING AND ARTIFICIAL VENTILATION

Natural spontaneous breathing increases the inner thorax volume by contracting the respiratory muscles. This creates a negative pressure in the lungs resulting in air being drawn in. However, artificial ventilation employs a reverse principle. The ventilator creates a negative pressure and thus pushes the respiratory gas into the lungs. The gas transport is called lung ventilation during both spontaneous and artificial ventilation.

The breathing apparatus model in the illustration shows both principles of ventilation as it would take place in a homogenous lung area. Roughly simplified, it shows an extensive analogy of ventilation during spontaneous breathing and artificial ventilation. However, the model insufficiently demonstrates the actual physical conditions. In fact, the lungs do not consist of only a single area, but are made up of several million microscopically small vesicular structures, known as the alveoli.

The realistic model demonstrates the difference between physiologic breathing and artificial ventilation. Every individual alveoli has different mechanical features and, depending on the position of each alveoli, the pressure of the ventilator or the respiratory muscle has a different effect: Breathing pressure primarily ventilates the upper lung areas while spontaneous breathing has a stronger effect on the lower areas close to the diaphragm.

The relevant difference between ventilation and breathing does not solely concern lung ventilation, but rather the pressures effecting the lungs also display discernible differences: During ventilation, the lungs are constantly exposed to positive pressure, even during expiration. During the expiration phase, the ventilator provides a low pressure, known as PEEP*. During spontaneous breathing, pressure in the lungs is temporarily lower than the ambient pressure. In addition, pressures arising in the lungs during artificial ventilation may be many times higher than the pressures applied to the lungs during physiologic breathing.

* PEEP = Positive End Expiratory Pressure
The breathing-related changes of the pulmonary pressures indicate that lung ventilation during artificial ventilation differs considerably from spontaneous breathing. The question of whether it is inferior has been an object of long-standing debate ever since the first implementation of mechanical ventilators. This discussion is not only limited to the effects on the lung as an organ and its ventilation. The effects on other organs are of concern as well. The cardiovascular system is of special importance, since it is more affected during ventilation than it would be during breathing.
MISINTERPRETATION OF SPONTANEOUS BREATHING AS AN INTERFERENCE FACTOR

An exceptional change in dealing with the patient’s spontaneous breathing is currently taking place in the development of artificial ventilation. In the early years, keeping up the vital gas exchange function during abnormal breathing conditions stood at the center of attention.

In those earlier days, mechanical ventilation was a measure in which the focus was on the lung. Potential negative effects to the lungs and other organs were accepted, if they were understood at all. The technical means for detecting and preventing side effects were not available at that time. The technical inadequacy of the ventilators at that time was even more serious: Spontaneous breathing often resulted in an impaired device functionality and, consequently, had to be suppressed by medication.

Since this course of treatment with a smooth transition from artificial ventilation to natural breathing was not possible then. The devices of that time only enabled the abrupt transfer from artificial ventilation to spontaneous breathing. Mixed forms of machine ventilation and spontaneous breathing did not exist.

If chemical paralysis of the patient using muscle relaxants is considered a questionable measure with today’s knowledge, motivations of the past treatment concepts still remain current today. The calm and immobilized patient was misinterpreted then, unfortunately also frequently still today, as a patient without complications. These complications were recognized only once they had already become untreatable and a fatal ending of the patient had become inevitable.
An additional serious problem occurred, especially if chemical paralysis was applied as a routine measure: This routine sedation impeded the patient's preparation for the time after artificial ventilation and thus delayed weaning from mechanical ventilation (10). The patient's personal contribution in this pre-weaning phase in the form of spontaneous breathing is a mandatory requirement.

Ventilation with the Spiromat
Suppression of spontaneous breathing is necessary for disturbance-free mechanical ventilation. A smooth transition to natural spontaneous breathing is impossible.
THE DEVELOPMENT OF SPONTANEOUS BREATHING AS TREATMENT GOAL

One of the first attempts to prepare the patient for the time after artificial ventilation was undertaken in the middle of the last century. The effort necessary for the patient to trigger the mechanical breaths was increased. The intention was to train the patient by continuously increasing the respiratory effort for triggering a breath (no s). The trigger threshold was increased for this purpose. The goal was to prepare the patient for spontaneous breathing.

As soon as the patient was able to exceed a certain trigger threshold, the ventilation was abruptly switched over to pure spontaneous breathing. This kind of procedure has multiple disadvantages: The trigger effort burdens the breathing apparatus in a non-physiological manner and contributes nothing to lung ventilation.

One mode that allows for continuous transition from mechanical ventilation to spontaneous breathing to some extent was developed in the 1970’s and called SIMV* (18). This procedure enabled a spontaneous breathing for the first time during the mechanical expiration phase. This was applied then and still is being used today, especially during weaning. It’s acceptance seems to be decreasing, since SIMV in its original form has an essential disadvantage. The mechanical breaths may still be triggered by the patient, but spontaneous breathing is not possible during these strokes. This procedure thus only enables a ventilation pattern in which the mechanical breaths and spontaneous breathing take place successively. This resulted in phases where the patient is unable to breathe spontaneously.

There are no medical indications for such interruptions of spontaneous breathing. There are no technical reasons - at least not in modern ventilators.

The technical requirements for uninterrupted spontaneous breathing during artificial ventilation, however, were not found in previous years. It was not until the 1980’s that this technology became available for the first time. This point in time is considered a development milestone by many users. Conventional ventilation that did not allow spontaneous breathing at any time due to technical limitations developed into modern ventilation. A new ventilation mode with electromagnetic valves and a microprocessor controller met the requirements of allowing spontaneous breathing during artificial ventilation at any time.

* SIMV = Synchronized Intermittent Mandatory Ventilation
Ventilation with the Evita 2
The ventilation curves in the background demonstrate the development of the mechanic procedures. Bottom: controlled ventilation without spontaneous breathing, top: mechanical ventilation with uninterrupted spontaneous breathing.
The technical principle of respiratory gas delivery

**SPONTANEOUS BREATHING WITH CONVENTIONAL ARTIFICIAL VENTILATION**

In the diagram, valve control during inspiratory pauses of conventional ventilation is simplified for a device without showing the additional function „base flow“. This function will be explained later. The first step is to describe a ventilation with parameters set by the operator and without the capability of adapting the ventilation pattern by the patient. This is controlled ventilation.

The inspiratory valve opens during the inspiratory phase until the respiratory gas of the respective breath is completely delivered. Longer mechanical inspiratory times are followed by a phase during which no respiratory gas flows until the end of the inspiratory phase. During the entire mechanical inspiratory time, the expiration valve is completely closed during controlled ventilation. The expiration valve opens at the beginning of the expiratory phase and releases the respiratory gas. During controlled ventilation, the inspiratory valve is closed during this phase. Only a timer is required to achieve inspiratory/expiratory phase control. These are the essential functional components of the ventilator for controlled ventilation.

If the ventilator must adapt to a breathing patient, the technical challenge becomes more complex. In this case, both valves need to be able to react to the spontaneous patient breathing. These valves must be ready to open at any time.

This principle was first realized in the expiration phase. Spontaneous breathing in the mechanical expiration phase has been part of ventilation technology standard since the 1970’s. To implement this into the mechanical inspiratory phase proved a little more difficult. Special ventilation modes enabled the patient to inhale during the mechanical inspiratory phase and thus to inhale more respiratory gas. But it was impossible to exhale in the mechanical inspiratory phase and thus the patient could not exhale before the mechanical expiration phase. The ventilator maintained the expiration valve closed during this phase and kept the patient from exhaling.

The first ventilator to allow an uninterrupted spontaneous breathing at any time during mechanical ventilation was Evita in 1988 using the BIPAP/PCV+* mode.
It took more than a decade until another ventilator with this performance feature became commercially available. By now in modern ventilation, it is understood that a patient may breathe spontaneously at any time over the full cycle.

There are quality differences in respiratory gas delivery for spontaneous breathing that can be explained when taking a closer look at the technical devices for respiratory gas dosing. The quality of spontaneous breathing depends not only on the features of these gas dosing devices, but also on the interaction of the individual systems.
INSPIRATORY VALVES

For reasons of simplicity, the device components for dosing the breathing gas were previously described as „the inspiratory valve“ and „the expiration valve“. In fact, complex designs are necessary for gas dosage.

Since the beginning of mechanical ventilation in the early years of the past century, valve systems in conjunction with pressurized gas sources were used for dosing respiratory gas. Designs using blowers to supply respiratory gas to valve systems have existed since the middle of the last century. If the blower and the patient are separated by a membrane, then it becomes bellows ventilation as commonly used in anesthesia. Respiratory gas dosage via controlled blowers was introduced only recently.

Modern intensive ventilation uses two systems for respiratory gas dosage:
Proportional valves (so called flow valves) for flow control and controlled blowers.

Similar to a water taps, proportional valves can generate any given flow between 0 and a maximum value. Different from a water tap, these changes do not require seconds, but occur within milliseconds. The precision and dynamics required by a sufficient respiratory gas dosage exceeds the performance of the pneumatic valve control used in the past. This is the reason why modern ventilators commonly use electromagnetic valve drives.

For comparative purposes, an electromagnetic valve controls function like loud speakers, employing a live coil moving in a magnetic field. In a loud speaker, the coil deflection is performed depending on the current and it sets a membrane in motion. A tappet in an electromagnetically controlled valve is moved in a similar fashion. The movement must closely follow the current. If it does not, the music originating from the loud speaker will sound odd. In proportional valves, the gas flow will not be sufficient for controlling the gas dosage with the required accuracy.
The ventilator proportional valves must resist high mechanical stresses. These valves must also be maintenance free and able to function with oxygen in high concentrations. That’s why only high quality materials are employed: For example, the Evita’s proportional valve closing mechanism consists of a sapphire ring and a ruby ball.

The term „proportional“ explains the technical control principle of the inspiratory valve. The technical principle is of lesser importance in comparison with the ventilation quality resulting from it. The term „demand valve“ provides a good explanation: The valve must deliver exactly what the patient demands. If the patient demands a large amount, it will deliver a large amount. If the patient demands nothing, the valve will remain closed. Alternative designs exist with valves that never fully close. These deliver a small gas flow even if the patient does not require any respiratory gas. The gas will not enter the patient’s lungs but exits via an expiration valve. Such designs are called „demand valve with base flow“.
EXPIRATION VALVES

Requirements for the expiration valve are different, but not less than those for the inspiratory valves. It should allow the respiratory gas to flow freely at the beginning of expiration phase and must not pose a large resistance. But it must also be able to close quickly, e.g. in order to sustain a constant residual pressure in the airways. This pressure must then be maintained as constant as possible - even if pressure fluctuations occur in the airways due to spontaneous breathing activity. The expiration valve must be very sensitive and react very quickly.

These high requirements are met by membrane valves. As with most inspiratory valves, the drive mechanism of diaphragm valves is usually electro-magnetic. Differing from inspiratory valves, the shutter mechanism and drive are spatially separated and not contained in a common housing. Dräger ventilators thus use two different drive systems: Older designs contain an air path between the valve membrane and drive. There the membrane is moved pneumatically. Newer designs like the Evita Infinity V500* use directly driven expiration valves.

Hygienic aspects and issues concerning the device treatment create special requirements. The expiration valve comes into direct contact with the patient's exhaled air and thus should be autoclavable. The expiration valve must meet two requirements in respect to the daily device preparations: One, it should be light and removable by hand, so that cleaning procedure may be performed quickly. Secondly, it should also be robust and housed in a compact housing without other functional components such as sensors, so that no damage may be caused by the daily device usage.
As described before, the expiration valve is part of the patient system. Thus it should not only be evaluated as a functional component in the ventilator control, but also under logistical aspects during ventilator device operation. If the operator should decide not to disinfect the system, the circuit and the expiration valve should be offered as disposable components.

Expiration valve of the Evita Infinity V500
The valve mechanism is contained in a compact housing. The expiration valve may be exchanged manually without using tools. The illustration shows a disposable valve.
RADIAL COMPRESSOR

The technical principal of respiratory gas dosage via controlled blowers (compressors) differs from the proportional valves described so far. As the description „proportional“ does in the above context, the term „blower“ only describes the technical function. „Compressor“ describes the function during ventilation more accurately: Respiratory gas is compressed and administered to the patient in a dosed manner.

Two different types of compressors are used during ventilation: The radial compressor and the side channel compressor. Their common feature is the respiratory gas compression using centrifugal forces. This presents the decisive advantage that the respiratory gas may be taken from the ambient air or a low pressure source (like an oxygen concentrator). Ventilation may thus be performed without a central gas supply if required. The compressors differ in breathing gas flow and pressure control.

Radial compressors work like a type of carousel for gas molecules. A special wheel accelerates the gas and propels it outwards radially using centrifugal force - hence the name. Thus pressure builds up behind the wheel. Once the path is free, the gas flows away from the compressor and into the patient's lungs. Pressure and flow are controlled via the wheel velocity.

The maximum velocity of a radial compressor in a ventilator is 80,000 rotations per minute and is thus comparable to aircraft turbines. The radial compressor must be able to accelerate to high speeds within milliseconds. It must also operate at low noise levels and create low heat emissions. Knowing this, the development efforts for such a system may be easily understood.

Dräger uses radial compressors in various areas of ventilation. They operate Carina for intensive ventilation - a ventilator that was primarily developed for non-invasive ventilation, e.g. via breathing masks.
Radial compressors use their high speed to create nearly all pressure required for ventilation purposes without additional equipment. But in order to achieve pressure changes, they must alter their speed. Acceleration and deceleration of the compressor wheel should be performed without delay. In the high speed range, this is technically difficult and only achievable to a certain extent.

If almost completely delay-free pressure increases without quick speed changes are required, a side channel compressor in combination with an auxiliary valve offers an alternative in the gas dosing system of a ventilator.
SIDE CHANNEL COMPRESSOR WITH AUXILIARY CIRCUITRY

Side channel compressors use a less known and rarely used principle for gas compression. A special type of wheel rotates in a chamber and sustains a small amount of gas in a helical rotation. It is positioned to draw in ambient air into the chamber and propel it into the connected side chamber, the so called side channel - hence the name. The respiratory gas then has a slightly increased pressure. An auxiliary circuit is used to feed it to the patient if required. This auxiliary circuit consists of a valve and a compressor control and provides an adequate subsequent gas delivery into the side channel.

This gas dosage principle resembles the valve system already described. However, the respiratory gas does not originate from a high pressure source - it is delivered from an upstream blower. The side channel compressor with auxiliary valve is used in the Savina ventilator.

Side channel compressors with an auxiliary valve offer similar advantages for ventilation as the proportional valves: These compressors are able to generate very fast pressure changes, since they already store pressurized respiratory gas. The speed of the pressure change depends largely only on the auxiliary circuitry that is able to react extremely quickly if a gas dosage change is required.

The comparison of both compressors with the proportional valves described above shows specific advantages for both the compressors and the proportional valves. Compressors operate with ambient air and require no pressurized gas supply, these compressors are independent of a central gas supply. Proportional valves are able to deliver every desired respiratory gas flow and, as opposed to compressors, they are able to generate a constant respiratory gas flow in any required intensity. Both compressors and proportional valves have an excellent reaction during spontaneous breathing. But proportional valves require higher technological effort to achieve this performance.
The quality of spontaneous breathing depends not only on dosing systems and expiration valves in particular. The more important question is how well the two systems have been matched to each other.

The importance of why an optimal interaction of the individual components in the respiratory gas control will be described in the following chapter. The description on this interaction uses the term „inspiratory valves“ for reasons of simplification although it also applies to compressors.
CONTROL OF RESPIRATORY GAS DELIVERY DURING MECHANICAL VENTILATION

In artificial ventilation, the patient and ventilator are connected via the patient system. Every respiratory gas delivery by the inspiratory valves thus immediately affects the patient’s lungs. This especially applies to closed systems. For these types of systems, during the mechanical inspiratory phase, there is no other way for the respiratory gas than the way into the patient’s lungs. This at least applies for normal inspiratory pressure. Only dangerously high pressures trigger an additional safety valve that will open an external passage for the respiratory gas.

The patient’s total respiratory gas demand is unknown at the beginning of a breath. Neither the lung mechanics nor the intensity of a spontaneous effort is known before the triggering of a breath. Thus the ventilator must recognize the gas demand during the respiratory gas delivery and must continuously adjust the respiratory gas flow during the respiratory gas delivery. Adjusting the respiratory gas delivery is a balancing act, since the device must react without delay and yet without delivering too much gas. In a closed system, this would probably cause an excessive pressure in the lungs.

There are various possibilities for protective mechanisms against potentially dangerous pressures due to excessive respiratory gas flows. Closed systems differs fundamentally from an open system. The protection principle in a closed system ensures a control of the inspiratory valve. The inspiration valve’s performance is reduced for this purpose. It slowly opens and to an extent approaches the peak flow carefully. Thus the respiratory gas is delivered more slowly and the technically possible maximum valve performance is not completely utilized.

Dräger designed the open system as an alternative to the closed system. This system continuously controls not only the inspiratory valve, but also the expiration valve. If an excessive gas delivery occurs in an open system, the expiration valve may already be opened during the mechanical inspiratory phase. Through this the expiration valve contributes in the effort to avoid the generation of dangerous pressures. Thus the maximum performance of the inspiratory valve is completely utilized in this system.
The open system was first implemented by Dräger at the end of the 1970’s. It has been continuously further developed and since then has been available in all Dräger intensive care ventilators. Other suppliers followed on this path considerably later, and the open system with a controlled expiration valve slowly established itself in the ventilation technology.

There is a simple explanation for the slower than expected development: Many older ventilators use expiration valves that were simply overstrained with the requirements of the open system. Their reaction behavior was too slow. This forced some manufacturers to replace the expiration valve by a new design before they could dare approach the new control technology.
Spontaneous breathing in new ventilation modes

SPONTANEOUS BREATHING DURING MECHANICAL VENTILATION

The problem of conventional ventilation is that it offers only mechanical ventilation breaths that do not allow spontaneous breathing. If spontaneous breathing attempts occur in a closed system in spite of this, the disruptions described above will result.

If the patient attempts to exhale, this will increase system pressure. A patient’s sudden cough will have even more drastic effects. This will quickly result in the occurrence of pressures that will trigger an alarm. In addition, safety mechanisms may become active by aborting a mechanical breath or opening the system via a safety valve. Such disorders due to spontaneous breathing activities of older ventilators are due to technical factors.

The consequences of hampered or delayed spontaneous breathing are serious. Patients may feel stressed if they cannot exhale during the mechanical breaths. A continuous hampering of spontaneous breathing will result in stress. If a forced expiration during a mechanical breaths triggers an alarm due to the high airway pressure, this may cause additional stress. If the pressure increases further, the interrupted mechanical breaths will result in reduced ventilation and the opening of the safety valve may result in a temporary PEEP loss.
In contrast, very good reasons exist for allowing spontaneous breathing at any time. The continuous availability of spontaneous breathing may possibly be perceived as less stress for many patients. This will increase the readiness to do additional respiratory work that may, for example, accelerate the weaning process. The patient will receive more “respiratory freedom”.

Furthermore, unhampered spontaneous breathing may enable optimization of analgesic sedation. Implementation of such measures may become unavoidable in many cases, but at least there is an indication that may become obsolete if an adequate ventilator is employed: The use of sedatives solely for suppressing spontaneous breathing (56).
SPONTANEOUS BREATHING WITH BIPAP/PCV+ AND APRV

A solution to the clinical problems caused by conventional ventilation was achieved in the 1980’s via two different approaches in Austria and the USA. All the more remarkable is that neither of the two workgroups was intent on achieving spontaneous breathing during mechanical ventilation. Rather, both groups were looking for the solution to a specific clinical problem and in this way achieved a new viewpoint regarding spontaneous breathing during mechanical ventilation.

The Austrian workgroup was searching to improve weaning, created the „universal weaning process“ and called it BIPAP/PCV+(6). The American workgroup was looking for ventilation improvements, discovered an effective carbon dioxide expiration by short-term pressure relief and called it APRV (19). Both procedures are largely identical in terms of the control behavior. In respect to their operation, they differ only in various parameters and in their clinical field of application.

The historic contribution of having made the decisive advance in conventional ventilation must be awarded to both groups in equal shares. But it took years for the clinical benefit of both modes to be recognized. The continuous availability of spontaneous breathing during mechanical ventilation slowly formed terms used internationally, such as „Room to Breathe“.

* APRV = Airway Pressure Release Ventilation
BIPAP/PCV+ and APRV
Top: The BIPAP/PCV+ mode is represented as an addition of pressure controlled ventilation and spontaneous breathing.
Bottom: APRV – Short term pressure relief from a higher airway pressure.
Both BIPAP/PCV+ and APRV are modes whose ventilation is adjusted by choosing the desired ventilation pressure. Such modes are termed “pressure controlled”. In contrast to these, modes that require the selection of volumes are called “volume controlled”. „Room to Breathe“ became available to volume controlled modes only around a decade after it was introduced to the pressure controlled ventilation (BIPAP/PCV+).

In contrast to pressure controlled ventilation, Room to Breathe was not integrated into volume controlled ventilation by using new modes, but by using an additional adjunct in already established modes, such as SIMV. Dräger introduced this feature under the name „AutoFlow“. Since then it is available optionally for the total spectrum of volume controlled modes. With respect to the „Room to Breathe“ concept, other suppliers restrict themselves to special modes or offer this feature exclusively for pressure controlled ventilation.

The combination of „Room to breathe“ with a volume controlled mode featuring a variable mechanical ventilation proportion was especially promising. A mode with a variable number of mechanical breaths that change depending on spontaneous breathing, as opposed to SIMV with a fixed number of breaths, has been known for some time under the name of MMV*. When the MMV mode is combined with the AutoFlow option, spontaneous breathing receives a synergistic benefit.

In addition to the previously described benefit of spontaneous breathing during mechanical breaths, another advantage becomes obvious in MMV: The frequency of mechanically delivered (mandatory) breaths is dependant upon spontaneous breathing. The more a patient breathes spontaneously, the lower the frequency of mandatory breaths. The result was a rapid, wean from the ventilator especially in short term ventilation may be achieved. This applies, for example, to treatment after anesthesia.

* MMV = Mandatory Minute Ventilation
Weaning using the MMV mode may be performed on uncomplicated cases, e.g. in post-operative ventilation, during the entire process without changing the ventilation mode (15). Once the spontaneous breathing satisfies the entire demand during MMV-AutoFlow, the number of mechanical breaths is reduced to zero. Such a control principle will lead to the weaning goal in a harmonizing and continuous fashion.

In addition to the adjustment of the ventilation to active spontaneous breathing described herein, another possibility exists for improving treatment quality: Through adjusting the ventilation to the passive features of the breathing apparatus, harmful side effects may also be reduced. Historically, such efforts are older than the development of free breathing capability at the end of the 1880’s. Adjustment of ventilation to the passive features began in the 1950’s.
Passive features of the breathing apparatus

COMPLIANCE AND RESISTANCE

Orientation for adjusting ventilation parameters was formerly provided by standard adjustments. Adjustments to the device were individually adapted to the patient only after knowledge of the breathing apparatus features were accumulated. The passive breathing apparatus features may be described by static pressure-volume relationship and with the behavior of flowing gases.

The static pressure-volume relationship serves to illustrate the elastic features of the breathing apparatus. Volume change due to a defined pressure change is described as elasticity. Compliance is a measure for the elasticity. In simplier terms, it may be demonstrated like a balloon that undergoes a gradual pressure increase while the balloon volume is monitored.

At the beginning of the experiment while the balloon is barely filled, a type of threshold must be overcome. In the initial stage, the balloon will fill only slowly while the pressure is increased continuously. Once the balloon has been only partially filled, further pressure increases will result in considerable volume increases. This threshold is called opening pressure. The extension of the breathing apparatus proceeds comparitively: An opening pressure also exists there, combined with an overlying area in which pressure increases result in considerable volume increases. Continued expansion of the breathing apparatus will then result in smaller volume increases. However, it is not possible to explain this phenomenon with this model. The breathing apparatus becomes increasingly more rigid with higher expansion.

Apparently the breathing apparatus has an optimum range in which little pressure change will effect large volume changes and there are threshold areas above and below it.

In addition to the static pressure-volume relationships, the behavior of flowing gases may provide additional knowledge of the mechanics of the breathing apparatus. A driving pressure delta must exist in order for gases to flow. For example, if the respiratory gas should flow from the oral cavity into the alveoli, there must be a pressure difference between the pressure in the upper airways and alveolar pressure. Therefore, the strength of the respiratory gas flow is not only dependent
on the pressure difference, but also on the airway resistance. This phenomenon can also be illustrated by a simple experiment: When brewing coffee, the coffee filter resists the flowing coffee to a certain extent. If two coffee filters are used, the coffee flows even slower due to the higher resistance.

This simple experiment provides conclusions regarding ventilation: If the resistance in the airways increases, a larger pressure must be applied in order to achieve the same respiratory gas flow. The resistance of airway flow are quantitatively referred to as resistance.

A summary view of compliance and resistance establishes the following with respect to respiratory mechanics: Compliance determines how much respiratory gas will eventually be transported into the lung and resistance determines how quickly this proceeds.
RISK FACTORS DURING ARTIFICIAL VENTILATION

Ventilation inevitability effects changes in the respiratory mechanics and thus requires continuous adaptation. Pressure limitation is a simple adjustment during volume controlled ventilation and has been used since the middle of the last century during long term ventilation.

During volume controlled ventilation, the delivered volumes remain constant even when lung mechanics change. In the original form, breath volumes, respiratory gas flow, the time pattern were predefined.

Ventilation with constant tidal volumes and constant flow may effect local mechanical stresses if the respiratory gas spreads unevenly in the lungs. Such disruptions in gas distribution are especially characterized for diseased lungs. This can be illustrated with a simple lung model, the two compartment model. Both compartments differ in their resistance.

The model illustrates how increased airway resistance in individual lung areas may impede gas distribution. If a tidal volume with constant flow is delivered under these circumstances, the compartment with the lower resistance is preferentially ventilated.

The effects of different ventilator modes result in different mechanical stresses on the lung tissue. The compartment with the smaller resistance may be temporarily overstretched. Pressure differences may arise between the compartments that may result in gas transport from one compartment into the other and thus effect so-called Pendelluft breathing. Mechanical stresses due to pressure differences may cause local tissue damage. If this Pendelluft phenomena continues over a period of time, a serious complication during ventilation affecting the entire organ may occur systemically.

Lung parenchma regions areas with little resistance and high compliance are especially vulnerable. They are able to take in high volumes in a short time, which could possibly possibly lead to overdistention. Unfortunately, these are often the healthy areas of the lung. The amount of healthy compartment may decrease rapidly during ventilation if to high volumes are applied. This means that ventilation with volumes that are too high endangers the healthy areas and may reduce their proportion of health vs diseased lung tissue. Ventilation with inadequate pressures, volumes and flows may cause the opposite of the desired therapeutic goal - thereby creating ventilator induced lung injury.
Not only high pressures are problematic. During the expiration phase, pressures that are also bear risks. A PEEP value which is too low can have the effect that lung areas will open and close during each breathing cycle. This will expose them to cyclical strains.

The damages listed up to now were previously distinguished due to their primary cause. Thus, an overextension due to volume was called volutrauma and damage due to pressure was called barotrauma. The effects of a PEEP value which are too low were called atelectrauma. In recent times, this differentiation was abandoned and damages to the lungs caused by ventilation are now called VILI (Ventilator Induced Lung Injury).
LUNG PROTECTIVE VENTILATION - MANUAL ADJUSTMENTS

There are various ways to avoid unfavorable volumes and pressures during ventilation. When ventilating a damaged lung, conventional knowledge states that a tidal volume of 6 milliliters per kg ideal bodyweight and ventilation pressures of thirty mbar/cmH$_2$O should not be exceeded without reason (3). The PEEP is usually higher than five mbar/cmH$_2$O. Special disease patterns may require ventilation pressures and volumes that deviate considerably from the values listed above. In rare and extreme cases, PEEP values of up to twenty mbar/cmH$_2$O are necessary in order to enable gas exchange.

The selection of individual suitable pressure and volume parameters is always a balancing act between optimum gas exchange and harmful side effects. There are efforts to minimize the side effects which may be caused by constant flow. An example illustrating this effort would be the ventilator delivers respiratory gas at a constant flow rate only at the beginning of a breath and then continues to gradually reduce the flow. This is known as decelerating flow.

Such flow profiles may be adjusted directly on some ventilators. This may also occur as a consequence of the user performing additional adjustments on the ventilatory pressure: If the flow is limited to a maximum value, the ventilator is able to deliver a high flow only at the beginning of the breath. It must then reduce the respiratory gas delivery in order to avoid exceeding the adjusted pressure limit.

The pressure limit is adjusted using the additional parameter $P_{\text{max}}$. It is usually selected so that the tidal volume will still be completely delivered while the mode remains to be volume controlled. Therefore, the time for respiratory gas delivery is dependant on the mechanical properties of the lungs. The higher the resistance, the longer the gas delivery will take. The temporal variation of decelerating flow and pressure limitation is shown in the illustration.
Pressure limited ventilation with decelerating flow causes effects that may in turn be explained using the two-compartment model. In contrast to constant flow ventilation, ideally both compartments are ventilated similarly. The pressure difference between the two compartments is then smaller. The more-compliant compartment fills somewhat readily, but no Pendelluft breathing will occur. If this model is transferred to the lungs, the advantage is as follows: Both high resistance compartments are appropriately ventilated without overdistending the compartments with low resistance.

The essential advantage of pressure limited ventilation is the continuous flow adjustment during resistance changes. The higher the resistance, the slower the flow will decelerate - the respiratory gas delivery is distributed over a larger time period. The disadvantage of a manually adjusted pressure limitation is that the clinician must constantly check if the pressure limitation is adequate. Under certain circumstances, the tidal volume could be delivered with a reduced ventilation pressure and the pressure limit would then have to be adjusted manually.
If the compliance changes in the process of volume controlled ventilation, it will affect the ventilation pressure. A stiff lung requires a higher pressure for the same volume and vice versa: If the compliance increases, respiratory gas may be delivered with reduced pressure. The smaller illustration on the next page, top left, shows the variation of pressure and increasing compliance during pressure limited ventilation.

Remarkably, the pressure does not change at the beginning of the breath. Actually, an increasing compliance should result in the tidal volume being delivered with reduced pressure. Actually, a compliance increase in the pressure limited ventilation has no effect on the peak pressure. Only after the tidal volume has been completely delivered will the pressure decrease to a smaller value. If the peak pressure should also be reduced to this value at the beginning of the breath, the clinician must make a manual intervention. A reduction of pressure limit is required in order to extend the gas delivery over a longer period of time during the breath. After compliance changes, this adjustment must be repeated continuously. In pressure limited volume controlled ventilation, no automatic control has been implemented for this.

Automatic pressure adjustment in volume controlled ventilation has been known since the 1990’s. Dräger introduced it with the AutoFlow option in Evita 4 - an option that has already been described relating to spontaneous breathing.

In summary, the AutoFlow option offers two essential features: It allows spontaneous breathing during volume controlled ventilation at any time and automatically adjusts the pressure. Pressure control during AutoFlow is shown in the large illustration on the following page.
In AutoFlow, the actual delivered volume is compared with the adjusted tidal volume at every mechanical or spontaneous breath. If the delivered value was too high, pressure is automatically reduced by up to three mbar/cmH$_2$O in the next ventilation cycle. This will then result in a lower volume. However, if too little volume was delivered, ventilation pressure in the next ventilation cycle will increase, resulting in a volume increase. As in conventional volume controlled modes, a limit protecting against pressure values that are too high may be entered in AutoFlow. The pressure cannot increase from this point onward.

The Babylog 8000 plus for premature infant ventilation introduced a mode essentially the same as AutoFlow. There it is called guaranteed volume pressure controlled ventilation with the designation VG*. Other manufacturers introduced automatic pressure control into volume controlled ventilation, without combining it with spontaneous breathing that is available at any time.

* VG = Volume guarantee
Systematics and control of the new modes

THE NOMENCLATURE PROBLEM

In the previous chapters, the modes BIPAP/PCV+, APRV, SIMV and MMV were presented as four examples of ventilation modes. AutoFlow and VG were described concerning additional ventilation mode adjuncts. The goal of the following remarks is first, to give an overview of the most established modes and adjuncts, and to provide some orientation into the nomenclature jungle. A deeper insight into the nomenclature systematic is provided by a separate Dräger series booklet with the title „Ventilation modes in the ICU“.

There is a fairly confusing multitude of designations for modes and their supplemental adjuncts. At first the mode names provide no logical connection. They describe the individual modes, but give little reference to their classification into a system. In addition, manufacturers have mainly taken their separate paths concerning nomenclature, and these have not always been constructive. Previously, some companies have intentionally searched for their own mode designations in order to distinguish themselves from others. Also misleading was the fact that similar ventilation modes sometimes used different designations in different fields of application.

The confusion caused by conventional nomenclature was the major reason, why ventilator operators have been demanding a unified nomenclature over all areas of ventilation for years. This is especially a challenge for those manufacturers who offer ventilators for the various fields of acute medicine - such as intensive medicine, emergency medicine, or anesthesia. The nomenclature should thus be as logical and self-explanatory as possible. A clear and cross-functional unified nomenclature avoids misunderstandings and thus perhaps human error.
Classification of the modes into a system could be done by applying various criteria. If the criterion of respiratory effort is selected, then there are two categories: Mandatory modes and mechanically supported spontaneous breathing modes. With mandatory modes, the ventilator performs the ventilatory effort mainly independent of the patient’s respiratory activity. During mechanically supported spontaneous breathing modes, the patient and machine share the effort. Mechanical support thereby adapts itself to the patient’s respiratory activity leading to improved synchrony. The common feature of both categories is the trigger mechanism that is practically identical for both the mandatory modes and the mechanically supported current spontaneous breathing modes. Similarities exist in both categories concerning ventilation adjustment parameters.

The essential distinguishing feature between the mandatory modes and the mechanically supported spontaneous breathing modes is the difference in the way how the time length of the breath is set. In the mechanical modes, breath duration is adjusted by the clinician using time parameters such as Tinsp, I:E and f or RR*. In the spontaneous breathing modes, temporal stroke variation is no longer adjusted by parameters, rather results from the lung mechanics and the patient’s spontaneous breathing activity.

* $T_{\text{insp}} =$ inspiration time, $I:E =$ Inspiratory/Expiratory ratio, $f =$ respiratory frequency (ie: $RR =$ respiratory rate)
DESCRIPTIONS OF MODES AND FUNCTIONS

Further classification of the ventilation modes is performed based on other differentiation criteria: Within the mandatory modes, classification is done depending on the adjustment parameter that significantly determines lung ventilation. The result is then the two categories of pressure controlled and volume controlled modes already described above. This rough classification of the mandatory modes can be found in a designation prefix of the new nomenclature introduced by Dräger: PC for pressure controlled or VC for volume controlled modes. Mechanically supported spontaneous breathing modes are designated by the prefix SPN (spontaneous). In spontaneous breathing modes, ventilation is normally adjusted via a pressure parameter. That’s why this mode is not differentiated by the prefixes PC and VC.

The actual mode designation essentially describes the interaction between the patient and ventilator. For example, an important interaction is the triggering of mechanical breaths by the patient. In mechanical ventilation, we distinguish between the different phases within a cycle, in which the ventilator is ready for the triggering of a breath. This trigger behavior is a differentiating feature that is reproduced in the mode designation.

The name AC (Assist Control) thus describes a mode allowing the patient to trigger mandatory breaths at any time during the expiration phase. The SIMV mode, however, uses a breathing cycle that allows the triggering of a breath only at certain times at the end of the expiration phase. The name CMV (Continuous Mandatory Ventilation) describes a mode that does not enable patient triggered breaths. In this case, the respiratory rate must be user defined by using fixed numbers - the ventilation frequency.
The CPAP (Continuous Positive Airway Pressure) mode in its pure form uses neither mandatory breaths nor mechanical pressure support. Here, the ventilator delivers the respiratory gas without performing mechanical effort.

Designation extensions describe special control functions that are only briefly mentioned here and described in more detail in the booklet „Ventilation Modes“. Examples include the previously described functions AutoFlow and VG, as well as support functions with the designations PS (pressure support) and VS (volume support).

Exceptions within the spontaneous breathing modes are those using neither time parameters nor ventilation adjustment parameters. Here, both the time length and the depth of the ventilation are not adjusted by the operator. They are controlled by the patient’s respiratory drive. This mode includes PPS (Proportional Pressure Support), PAV (Proportional Assist Ventilation), NAVA (Neural Adjusted Ventilatory Assist).

<table>
<thead>
<tr>
<th>Prefix</th>
<th>Mode designation</th>
<th>Designation extension</th>
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<tbody>
<tr>
<td>PC</td>
<td>BIPAP/SIMV+</td>
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<tr>
<td></td>
<td>APRV</td>
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<td>PPS</td>
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Mode designation structure
The prefix describes the parameter effecting lung ventilation. The actual mode designation describes the interaction between device and patient. Additional designation extensions describe additional control functions.
CONTROL CONCEPT BIPAP/PCV+ AND APRV

Modern ventilation is characterized by spontaneous breathing being available at any time and automatic lung mechanics adaption. This was compared to the conventional modes represented by BIPAP/PCV+ and AutoFlow in the previous chapter. Here the patient can not always breathe spontaneously and an adaption to the lung mechanics requires manually performed measures. The following chapter now describes how ventilation parameters are adjusted on the patient and how modern mechanical ventilation is used in clinical application.

Compared with conventional modes, the introduction of the BIPAP/PCV+ mode opened up new paths of operation. Adjustment of the original Evita 1 in the 1988 was limited to only four parameters. In this case, the ventilation pressure was adjusted via the upper pressure level \( P_{\text{high}} \) and the lower pressure level \( P_{\text{low}} \). The mechanical inspiratory time was defined with the duration of the upper pressure level \( T_{\text{high}} \), while the mechanical expiration time was defined by the duration of the lower pressure level \( T_{\text{low}} \).

This represented a rather simple operating concept. The illustration reproduces the original BIPAP/PCV+ adjustment parameters in an Evita 1 in comparison with the conventional ventilation mode VC-SIMV.

The simple BIPAP/PCV+ operating concept with only four parameters only found limited acceptance. Some operators valued the simple principle with direct access to pressures and times and used the possibility to adjust extreme Inspiratory/Expiratory (I:E) time ratios. Others demanded the conventional adjustment they were used to from previous conventional modes.
The second generation of pressure controlled modes with free spontaneous breathing possibility fulfilled both requirements. The BIPAP/PCV+ operation was adapted to the conventional SIMV while the operating concept with four parameters was offered in the APRV mode.

BIPAP/PCV+ and APRV are nearly identical in respect to their behavior during spontaneous breathing and their reaction to changes in lung mechanics. They mainly differ in their operating concept and their previously described historical development. An additional difference is their clinical application.

The original BIPAP/PCV+ adjustment parameters were derived from an adjustment of a conventional SIMV. With pressure parameters, the BIPAP/PCV+ $P_{low}$ corresponds to the SIMV PEEP. The upper pressure value is directly adjusted as $P_{high}$ in the BIPAP/PCV+, in SIMV it is a result of adjusted tidal volume and lung mechanics and designated plateau pressure $P_{plat}$. In BIPAP/PCV+, the time values for inspiratory time were originally entered as $T_{high}$ and $T_{low}$. In SIMV they were a result of the adjustments of frequency (f) and inspiratory/expiratory ratio (I:E). In newer ventilators, the BIPAP/PCV+ and SIMV times are adjusted via the same parameters.
INTRODUCTION TO MODERN MODES

Ventilation with BIPAP/PCV+ and APRV may be performed at different times. It can be performed after a previous conventional VC-SIMV ventilation by changing the mode. Alternatively, the ventilation can begin from the start using BIPAP/PCV+ and APRV. Adjustment recommendations were published for both cases (32). The illustration reproduces the parameter adjustments in BIPAP/PCV+ ventilation.

When changing over from a volume controlled ventilation with conventional VC-SIMV, the time adjustment is transferred to BIPAP/PCV+ ventilation and remains unchanged. Thus, no new time parameters need to be adjusted since they are identical for conventional SIMV and BIPAP/PCV+. Before switching over to APRV, the values for $T_{\text{high}}$ and $T_{\text{low}}$ need to be converted from the SIMV adjustment values for breathing frequency and I:E ratio.

Regarding the pressure parameters, the lower pressure value in BIPAP/PCV+ is identical to the PEEP of the SIMV adjustment. During APRV however, the lower pressure value is adjusted via the parameter $P_{\text{low}}$. The upper pressure value needs to be readjusted for both BIPAP/PCV+ and APRV using parameters $P_{\text{insp}}$ during BIPAP/PCV+ and $P_{\text{high}}$ during APRV. The adjusted pressure value is dependant on the plateau pressure previously measured during VC-SIMV ventilation. After converting the mode, we recommend careful monitoring of the tidal volume for deviations from the target value, so that the ventilation pressure may be immediately readjusted.

If BIPAP/PCV+ or APRV are already implemented at the start of ventilation, clinical literature recommends a ventilation pressure of 12-15 mbar/cmH$_2$O above PEEP (32). Subsequently, the tidal volume should be observed and the ventilation pressure readjusted if required. Thereby, the breath volumes of lung protecting ventilation (3) should not be exceeded.
The velocity of respiratory gas delivered during BIPAP/PCV+ and APRV is adjusted via the parameter known as slope. In VC-SIMV, the respiratory gas delivery is controlled by the flow parameter. During the use of constant flow, there is no adjustment available for the selection of a pressure ramp.

Only few parameters need to be adjusted for the transition from a conventional VC-SIMV ventilation to VC-SIMV AutoFlow. In this case, the ventilator remains in volume controlled ventilation. Therefore, the clinician does not change the process, he only activates AutoFlow as an additional adjunct in the current mode. In AutoFlow, the parameter is not available as an adjustment for the respiratory gas velocity. It is adjusted via the pressure slope for both AutoFlow and BIPAP/PCV+.

**Initiating BIPAP/PCV+ ventilation**

1.) When switching over from a previous VC-SIMV ventilation, the plateau pressure serves as guideline for the adjustment of the ventilation pressure $P_{\text{insp}}$. The tidal volume $V_t$ is observed and altered by readjusting the ventilation pressure as required.

2.) At the start of BIPAP/PCV+ ventilation without previous conventional SIMV ventilation, we recommend an $P_{\text{insp}}$ adjustment of 12-15 mbar/cmH$_2$O with subsequent monitoring of $V_t$ and readjustment of $P_{\text{insp}}$ corresponding with 1.) as required.
PRESSURE LIMITED, AUTOFLOW AND BIPAP/PCV+ PUT INTO PRACTICE

A pressure limit during volume controlled ventilation already suffices to largely avoid potentially harmful pressures. During this mode, the application of an adequate pressure limitation may achieve an improved gas distribution in an inhomogeneous lung, due to the fact that instead of a constant flow, here a decelerating flow is generated. The advantages of pressure limitation in regard to lung protecting ventilation are opposed by disadvantages in clinical practice: Changes in the lung mechanics require the operator to manually readjust, and spontaneous breathing is not facilitated.

AutoFlow provides a remedy for both problems. During VC-SIMV, pressure and flow are controlled by this additional function and do not require readjustment by the operator. The target value of this control is the target tidal volume. The result is a ventilation with constant tidal volume and minimal peak pressure. Protection from an exceedingly high ventilating pressure is thereby provided by an additional security alarm system. As in other operations, it is activated through the high airway pressure alarm. All Dräger ventilators stop the delivery of breathing gas as soon as the airway pressure reaches a value 5 mbar/cmH₂O below the set high airway pressure alarm setting.

An essential advantage of AutoFlow in comparison with conventional ventilation appears during spontaneous breathing. The patient may breathe at any time. In practice, this means more comfort and less stress for the patient. During the mandatory breaths, the patient is no longer hindered from covering additional the demand for additional flow for spontaneous efforts. Essentially, breathing against becomes breathing with.

An alternative to volume controlled ventilation with AutoFlow is provided by the pressure controlled BIPAP/PCV+ mode. When compared with AutoFlow, BIPAP/PCV+ uses a different control behavior if there are changes in lung mechanics. In this case during BIPAP/PCV+ the ventilation pressure remains constant and the tidal volume varies.

The preferred fields of application of AutoFlow and BIPAP/PCV+ with their different control behavior are as follows: AutoFlow is always recommended when ventilation should be performed with constant tidal volumes and the ventilation pressure is
tolerable within certain limits. BIPAP/PCV+ is recommended when pressure-oriented ventilation is required. The delivered tidal volumes are not constant and may vary once the lung mechanics change.

In spite of its different control behavior during changing lung mechanics, BIPAP/PCV+ provides the same advantages as AutoFlow. It delivers respiratory gas with decelerated flow for improved respiratory gas distribution. It also offers the extraordinarily important clinical possibility to spontaneously breathe at any time. This enables a smooth transition from mechanical ventilation to spontaneous breathing - an advantage that becomes fully apparent during weaning. Spontaneous breathing during mechanical ventilation improves the gas distribution in the lungs and provides improvement in the patient’s ventilation especially in those lung areas that are only insufficiently ventilated by artificial ventilation (52). However, the advantages of spontaneous breathing are not limited to total lung ventilation; they also affect other gas exchange conditions.
Clinical application of the mechanical modes

CONDITIONS AND DISORDERS OF GAS EXCHANGE

Ventilation with its respiratory gas transport between atmosphere and alveoli is one requirement for pulmonary gas exchange. In addition, oxygen must constantly be transported via blood circulation from the alveoli and carbon dioxide removed. This sustains the respiratory gas diffusion in the alveoli. The lung circulation necessary for the respiratory gas transport is described as perfusion.

The gas exchange in the lungs must be viewed holistically in terms of ventilation and perfusion. The ratio of alveoli ventilation and lung capillary perfusion is a measure for the gas exchange conditions. It is designated as the ventilation / perfusion ratio and abbreviated $\frac{V}{Q}$.

The illustration shows the ideal conditions and possible causes for a reduced pulmonary gas exchange with an example of various alveoli. The middle alveoli shows the most favorable conditions for the gas exchange: Here the ventilation and circulation proceed undisturbed and the ventilation perfusion ratio is optimal. The left and right alveoli show the two forms of disorders within the lungs with their effects on the ventilation perfusion ratio.

Disorders of the pulmonary gas exchange are differentiated based on their causes of either poorly ventilated alveoli or insufficiently circulated alveoli. In the first case, alveoli may be disconnected from ventilation due to formation of atelectasis, so that no oxygenation of the blood takes place there in spite of good ventilation ($\frac{V}{Q}=0$). This type of phenomenon is called a shunt. In the second case, intact alveoli may be cut off from circulation e.g. by an embolism, so that no gas exchange takes place in spite of good ventilation. This case is designated alveolar dead space ventilation ($\frac{V}{Q}=\infty$). An additional requirement for the pulmonary gas exchange is an intact respiratory system. By acting as a type of „ventilation pump“, it ensures that the respiratory gas can even get to the gas exchange location.

Gas exchange disorders may be treated with targeted ventilation measures once the cause of the disorders has been identified. But in terms of mechanical ventilation, there is an additional reason for dealing with pulmonary gas exchange and especially with lung perfusion: Lung perfusion is significantly influenced by
ventilation itself. Spontaneous breathing influences lung perfusion as well. A comparison between ventilation and spontaneous breathing in terms of their effects on lung circulation will result in the same conclusions as a comparison in terms of lung ventilation. The natural way is the better way.

Natural breathing usually improves lung perfusion and artificial ventilation impedes it. When employing ventilation modes in clinical practice, the question is not only which targeted adaptations may be performed on the mode in order that a gas exchange disorder is reduced as much as possible. There is another challenge beyond this of not provoking such disorders by artificial ventilation.
BREATHING ADJUSTMENTS DURING OXYGENATION DISORDERS

Gas exchange disorders require fast reactions for corrective treatment. The risk is that the patient cannot absorb sufficient oxygen and dispense sufficient carbon dioxide by his own means. A respiratory insufficiency will then occur. Treatment depends on the present gas exchange malfunction. Basically, differentiation is made between two categories: Oxygenation disorders and alveolar ventilation disorders.

A differentiation of the gas exchange disorder is performed in the ICU via blood gas value measurements. If the arterial oxygen concentration is too low, this is considered as the leading symptom for an oxygenation disorder. This may be due to a change of the ventilation/perfusion ratio, e.g. due to a gas exchange area that is too small. Thereby, the volume at the end of the normal expiration, the so-called functional residual capacity (FRC), is reduced.

A reduction of the FRC is associated with an increased collapse of alveolar areas. The closure of the alveoli at the end of expiration and the reopening at the beginning of inspiration causes shear forces that may in turn cause fluid accumulations (edema) and even to lung tissue damage. This creates the conditions for gas exchange deteriorate further.

Thus an essential measure during oxygenation disorders is the restoration of an adequate gas exchange surface. In this case the goal is to keep the functional residual capacity stable and at an optimal value. This optimum provides not only the most favorable conditions for both the gas exchange and reduction of the respiratory effort. In addition, an increase of the fractional concentration of inspired oxygen (FiO₂).

As a guideline for the treatment of oxygenation disorders, an example for measures during BIPAP/PCV+ ventilation is described in the following (32): They are extended for AutoFlow and also shown in the illustration. The FRC increases due to a higher average airway pressure. The measures are:
Ventilation adjustments due to oxygenation malfunctions

In order to achieve an increase of the functional residual capacity (FRC), the average airway pressure may be increased using two different modes:

a) Similar increase of ventilation pressure and PEEP.

b) Increase of I:E ratio.

a) The similar changes of the upper and lower pressure values $P_{\text{high}}$ and $P_{\text{low}}$ in APRV or of PEEP and $P_{\text{insp}}$ in BIPAP/PCV+. During AutoFlow, only PEEP is increased.

b) The increased time of the upper pressure $T_{\text{high}}$ and reduction of the bottom pressure time $T_{\text{low}}$ in APRV, so that a higher I:E ratio is achieved. In BIPAP/PCV+ and AutoFlow, the I:E ratio is increased by adjusting $T_{\text{insp}}$ and $f$.

The selection of one of the two alternatives is dependent upon the individual case. In terms of the first possibility, clinical literature lists 30 mbar/cmH$_2$O as the limit for a reasonable ventilation pressure and a tidal volume of no more than 6 ml/kg body weight (3). In terms of the second possibility, the following effect needs to be considered: The expiration time may shorten so much that some lung areas can no longer be sufficiently ventilated. Thus, a remaining pressure above the selected PEEP will remain in the lung as intrinsic PEEP.
VENTILATION ADJUSTMENT DURING VENTILATION DISORDERS

Ventilation disorders are characterized by an increased arterial CO₂ concentration. Differing from the oxygenation disorders whose causes usually lie in the lung, ventilation malfunctions are mostly caused by a weakness of the respiratory system.

The reasons for reduced pump efficiency of the breathing apparatus are diverse and could be found in the respiratory control, respiratory muscles or in the airways. The central breath control could possibly be impaired e.g. by sedatives or a trauma of the brain.

During treatment of ventilation disorders, the ventilator takes over a proportion of the full pump efficiency of the respiratory system and thus guarantees alveolar ventilation. A possible adaption of the ventilation to the present disorder is shown in the illustration:

The measures for increasing the alveolar ventilation are possible as follows:

a) Increasing the ventilation pressure by increasing the upper pressure value $P_{\text{high}}$ in APRV or $P_{\text{insp}}$ in BIPAP/PCV+. The tidal volume is increased in AutoFlow.

b) The reduction of times $T_{\text{low}}$ and $T_{\text{high}}$ in APRV. In BIPAP/PCV+ and AutoFlow, the ventilation frequency is increased correspondingly.
Both measures intend to cause an increase of the respiratory minute volume and have specific advantages and disadvantages. If possible the increase of pressure and volume should not result in the overdistention of lung tissue.

More important than reacting to a gas exchange disorders by adapting the ventilation is to avoid these complications from the beginning. This involves careful use of medications that are capable of reducing the respiratory system performance with their side effects (56). These medications include sedatives that are paradoxically applied in order to improve the conditions for artificial ventilation.
EFFECTIVE GAS EXCHANGE AND SPONTANEOUS BREATHING THROUGH RECRUITMENT

If the lung condition deteriorates, the required measures for this are usually connected with higher stress levels for the patient. This further aggressive approach, in regards to ventilation is more encompassing than in regular usage. The issue at hand is not only the differentiation of measures in terms of how far excessive pressures affect the patient - e.g. concerning connection of the patient to the ventilator: This may be performed invasively via a tube or non-invasively via a mask. The matter of concern is stress in general. Increased ventilatory pressure in artificial ventilation is regarded as more invasive than the much smaller airway pressure during spontaneous breathing.

While there is no alternative for most measures with more aggressive tactics, seen from a clinical viewpoint few occasions give reason to suppress spontaneous breathing by sedatives. Ventilation modes that do not enable spontaneous breathing due to technical limitations and thus require suppression of returning spontaneous breathing are therefore problematic. They lose one of the few possibilities to reduce the invasiveness of a therapeutic measure.

Spontaneous breathing is possible in the modes BIPAP/PCV+, APRV and AutoFlow at any time and therefore additional natural breathing is possible. Due to contraction the diaphragm dome flattens and causes a ventilation of the lower sections of the lungs close to the diaphragm. Due to gravity exactly these areas are well circulated. However, the less circulated upper areas are less ventilated during spontaneous breathing and thus a balanced ventilation/perfusion ratio results for the entire lung. This provides good conditions for gas exchange for the entire lung.

The use of sedatives and especially relaxants deteriorates the conditions for gas exchange. Functional residual capacity is reduced and the gas exchange surface becomes smaller. The respiratory muscles weaken and artificial ventilation increasingly ventilates the upper and less well circulated lung sections. This results in an increasing formation of atelectasis in the lower lung regions.
Spontaneous breathing is partially able to remedy the negative effects of mechanical ventilation. Since spontaneous breathing strongly affects the basal-dorsal lung section, it thus reaches a type of „mechanical breathing problem zone“. If the alveoli in these sections are partly filled with fluid and collapsed, the spontaneous breathing may possibly reopen and reventilate them. This type of „revival“ of lung areas is also called „alveolar recruitment“.

Effects of spontaneous breathing on the gas exchange

The influencing factors classified regarding their effect on lung perfusion (top) and lung ventilation (bottom) and the general influencing factors (middle)
CONVENTIONAL WEANING AND UNIVERSAL WEANING MODE BIPAP/PCV+

The weaning process from the ventilator is a systematic reduction of the invasiveness in artificial ventilation. The goal is to guide the patient back to self-breathing, so that he/she will finally become independent of the ventilator. The weaning process is not a static process with a fixed schedule, it is a dynamic process.

During conventional ventilation, multiple weaning phases with a respective individual ventilation mode are differentiated. The left illustration shows a typical weaning process with the modes used in conventional ventilation. The following demonstrates how complicated the conventional weaning process can become.

In the course of the weaning process, the treatment team specifically reduces the proportion of mechanical breathing support, which the patient will then replace by natural respiratory effort. The mixed ventilation forms used in this case are also called augmented ventilation. Of these, two principles are used in conventional ventilation: On the one hand, there is intermittent ventilation and on the other, pressure supported ventilation.

During ventilation, e.g. SIMV, the mechanical breaths and spontaneous breathing are inter-mixed. The clinician thereby specifies the frequency and duration of the mechanical breaths. During pressure supported ventilation, however, frequency and duration of the mechanical support depend on the patient. The operator only defines the degree of the mechanical support.

Both intermittent ventilation and pressure supported ventilation provide smooth transitions to pure spontaneous breathing. In conventional weaning, the established practice is to use intermittent and pressure supported ventilation at the same time. In this case, e.g. in the SIMV mode, the spontaneous breathing between the mechanical strokes is supported with a mechanical pressure support.
The BIPAP/PCV+ weaning concept however uses only one mode over the whole process from beginning to completion. This it is also known by some clinicians as the universal weaning mode. The right illustration depicts a weaning course in BIPAP/PCV+ in the same phases as in conventional weaning.

The advantage of such a weaning concept is apparent: The operator neither needs to decide on whether to switch over between different modes nor deal with the combination of two modes. The patient benefits from this concept as well. The patient benefits not only from the improved gas exchange due to the patient's spontaneous ventilation, but also from the continuous weaning process.

Conventional weaning and with BIPAP/PCV+
Left: Different modes are used in conventional weaning:
1.) Pure mechanical ventilation without spontaneous breathing.
2.) Augmented intermittent: Spontaneous breathing and mechanical ventilation alternate.
3.) Augmented pressure supported: Mechanical support of spontaneous breathing.
4.) Pure spontaneous breathing without mechanical support.

Right: During weaning with BIPAP/PCV+, only one mode is used:
1.) BIPAP/PCV+ without spontaneous breathing.
2.) Spontaneous breathing only on the lower pressure level.
3.) Continuous spontaneous breathing on both pressure levels.
4.) Continuous spontaneous breathing - both pressure levels are adjusted to each other.
WEANING WITH BIPAP/PCV+

In the course of weaning, the invasiveness is gradually reduced and the inspired O₂ concentration is reduced to less than 50%. The I:E ratio is reduced so far that the expiration time takes longer than the inspiratory time.

Following the above adjustments, the PEEP may be gradually reduced to 7-9 mbar/cmH₂O in 2-3 mbar/cmH₂O increments. In the further course of conventional weaning, the ventilation mode would need to be changed to a mode allowing spontaneous breathing on the PEEP level. Weaning with BIPAP/PCV+ deviates from this scheme at a very decisive point: In BIPAP/PCV+, spontaneous breathing is possible from the start and is available at any time. The illustration shows an overview of the BIPAP/PCV+ weaning concept.

Weaning with BIPAP/PCV+ consists, like conventional weaning, of a gradual reduction of invasiveness. Various phases are recommended for the gradual reduction of the mechanical ventilation (32, 55):

1. Reduce inspiratory O₂ concentration to less than 50% (FiO₂ < 0.5).

2. Reduce Inspiratory/Expiratory ratio (I:E < 1).

3. Gradually reduce PEEP to 7-9 mbar/cmH₂O and reduce the upper pressure value (P_{insp}) adequately in order to lower the average airway pressure.

4. Gradually reduce the ventilation pressure amplitude until the pressure difference between PEEP and P_{insp} is 8-12 mbar/cmH₂O.

5. First reduce ventilation frequency to 8-9 strokes per minute. Then perform a further reduction of the I:E ratio and the frequency.

6. Gradually adjust both pressure values in steps to match the average airway pressure. Lower the PEEP value only after complete adjustment of the pressure values.
Weaning with BIPAP/PCV+
Explanations see text
SUCCESS FACTORS OF THE INDEPENDENT SPONTANEOUS BREATHING IN MECHANICAL VENTILATION

With increasing severity of a disease, mechanical ventilation often allows only little room for action. With increasing invasiveness of the ventilation, the danger increases of destroying what is actually supposed to be saved. In the end, the complications in ventilation often result in a „Circulus Vitiosus“ (vicious cycle). They result in a chain of adverse effects that allow fewer and fewer ways out of the situation.

In contrast to this, sometimes chains of events occur in ventilation that positively reinforce each other. An example for this is the free spontaneous breathing that is available at any time during mechanical ventilation.

Free spontaneous breathing allows the use of fewer sedatives. This reduces damping of the respiratory reflex. Over time, the strengthened spontaneous breathing will re-open atelectatic portions of the lung consequently increasing the lung's surface area for improvements in gas exchange. Henceforth, the ventilating pressures may be reduced since a part of the ventilation can now be provided by spontaneous ventilation. Lower ventilation pressure will facilitate spontaneous breathing during mechanical breaths.

The free spontaneous breathing in mechanical ventilation triggers a chain of events that positively reinforce each other and thus accelerates the therapeutic success.

The continually increasing complexity of equipment used to be a sign of progress in the past: More ventilation modes, more adjustment parameters, more additional functions. The BIPAP/PCV+ concept deviates from this principle. As BIPAP/PCV+ covers the full treatment spectrum from the beginning to the completion of weaning, it offers more flexibility than a a concept that is using multiple modes. The treatment team does not need to concentrate on the point in time for converting from one mode to the other. The measures can be dosed more finely over the course of the treatment and invasiveness can be precisely controlled. The continuous weaning process with BIPAP/PCV+ provides not only more flexibility, it also reduces the workload for clinic personnel.
Changing to a new ventilation mode during the weaning process requires careful defining of indications and preparation for the new mode. This is not applicable for BIPAP/PCV+. An additional workload reduction is provided by weaning with BIPAP/PCV+ and the reduced use of sedatives.

The BIPAP/PCV+ concept benefits hospital personnel with higher flexibility for therapeutic measures and work reduction within a continuous process. BIPAP/PCV+ is less a ventilation mode and more a ventilation therapy.
THE FUTURE OF SPONTANEOUS BREATHING IN INTENSIVE CARE VENTILATION

The introduction of the first ventilation mode allowing spontaneous breathing at any time took place in the 1980’s. Due to the development of precision valves with microprocessor control it became possible. At the same time, additional innovation such as graphical monitoring, were introduced to intensive care ventilation so that the eighties are now regarded as the beginning of modern ventilation.

For approximately 10 years, ventilation modes with spontaneous breathing being available at any time were only offered exclusively by one manufacturer. In the course of time, more and more clinicians have demanded this feature. It is now part of the standard equipment of all every intensive care ventilators, although they differ in quality.

Features that previously were exclusive top features of a ventilator are now standard equipment. They include various diagnostic functions, e.g. for evaluation of spontaneous breathing capability.

There is a tendency with intensive care ventilators parallel to universal devices to concentrate on certain essential performance features. This results in special devices being established with limited fields of application. The Carina ventilator is an example. This device Carina is highly suitable for mask ventilation. For other applications it only provides standard performance or does not include dispensable features. The Carina is a specialized and dedicated device for primarily non-invasive ventilation.
In the future, it will be possible to concentrate on specific performance requirements with additional specialized devices. An example of this is Savina 300, a ventilator with excellent performance in regards to spontaneous features and thus equivalent to the Evita Infinity V500 in this performance feature. However, in regards to many other performance features, the universal device remains superior to the specialized device: Complex equipment for monitoring lung functions, automatic ventilation control by knowledge-based systems, data management for the integration of ventilation into the acute medicine system will be reserved for the universal device for the foreseeable future.

In the future, additional specialized devices for intensive ventilation may possibly be developed, with one excelling performance feature and compromises in other areas. No compromises must be made in regards to a single feature in the future of intensive ventilation: Spontaneous breathing during mechanical ventilation is indispensable and must comply with the highest quality requirements.
Appendix

ADDITIONAL LITERATURE


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As of August 2015:
Dräger Medical GmbH changes to Drägerwerk AG & Co. KGaA.