We discuss the simple yet powerful ideas which have allowed to break the diffraction resolution limit of lens-based optical microscopy. The basic principles and standard implementations of STED (stimulated emission depletion) and RESOLFT (reversible saturable/switchable optical linear (fluorescence) transitions) microscopy are introduced, followed by selected highlights of recent advances, including MINFLUX (minimal photon fluxes) nanoscopy with molecule-size (~1 nm) resolution.

We are all familiar with the sayings “a picture is worth a thousand words” and “seeing is believing”. Not only do they apply to our daily lives, but certainly also to the natural sciences. Therefore, it is probably not by chance that the historical beginning of modern natural sciences very much coincides with the invention of light microscopy. With the light microscope mankind was able to see for the first time that every living being consists of cells as basic units of structure and function; bacteria were discovered with the light microscope, and also mitochondria as examples of subcellular organelles.

However, we learned in high school that the resolution of a light microscope is limited to about half the wavelength of the light [1–4], which typically amounts to about 200–350 nm. If we want to see details of smaller things, such as viruses for example, we have to resort to electron microscopy. Electron microscopy has achieved a much higher spatial resolution—tenfold, hundred-fold or even thousand-fold higher; in fact, down to the size of a single molecule. Therefore the question comes up: Why do we care for the light microscope and its spatial resolution, now that we have the electron microscope?

The first reason is that light microscopy is the only way in which we can look inside a living cell, or even living tissues, in three dimensions; it is minimally invasive. But, there is another reason. When we look into a cell, we are usually interested in a certain species of proteins or other biomolecules, and we have to make this species distinct from the rest—we have to “highlight” those proteins [5]. This is because, to light or to electrons, all the proteins look the same.

In light microscopy this “highlighting” is readily feasible by attaching a fluorescent molecule to the biomolecule of interest [6]. Importantly, a fluorescent molecule [7] has, among others, two fundamental states: a ground
state and an excited fluorescent state with higher energy. If we shine light of a suitable wavelength on it, for example green light, it can absorb a green photon so that the molecule is raised from its ground state to the excited state. Right afterwards the atoms of the molecule wiggle a bit—that is why the molecules have vibrational sub-states—but within a few nanoseconds, the molecule relaxes back to the ground state by emitting a fluorescence photon.

Because some of the energy of the absorbed (green) photon is lost in the wiggling of the atoms, the fluorescence photon is red-shifted in wavelength. This is actually very convenient, because we can now easily separate the fluorescence from the excitation light, the light with which the cell is illuminated. This shift in wavelength makes fluorescence microscopy extremely sensitive. In fact, it can be so sensitive that one can detect a single molecule, as has been discovered through the works of W. E. Moerner [8], of Michel Orrit [9] and their co-workers.

However, if a second molecule, a third molecule, a fourth molecule, a fifth molecule and so on are positioned closer together than about 200–350 nm, we cannot tell them apart, because they appear in the microscope as a single blur. Therefore, it is important to keep in mind that resolution is about telling features apart; it is about distinguishing them. Resolution must not be confused with sensitivity of detection, because it is about seeing different features as separate entities.

1.1 Breaking the Diffraction Barrier in the Far-field Fluorescence Microscope

Now it is easy to appreciate that a lot of information is lost if we look into a cell with a fluorescence microscope: anything that is below the scale of 200 nm appears blurred. Consequently, if one manages to come up with a focusing (far-field) fluorescence microscope which has a much higher spatial resolution, this would have a tremendous impact in the life sciences and beyond.

In a first step, we have to understand why the resolution of a conventional light-focusing microscope is limited. In simple terms it can be explained as follows. The most important element of a light microscope is the objective lens (Fig. 1.1). The role of this objective lens is simply to concentrate the light in space, to focus the light down to a point. However, because light propagates as a wave, it is not possible for the lens to concentrate the light in a single point. Rather the light will be diffracted, “smeared out” in the focal

**Fig. 1.1** Focusing of light by the microscope (objective) lens cannot occur more tightly than the diffraction (Abbe’s) limit. As a result, all molecules within this diffraction-limited region are illuminated together, emit virtually together, and cannot be told apart. Verdet [2], Abbe [1], Helmholtz [4], Rayleigh [3]
region, forming a spot of light which is—at minimum—about 200 nm wide and about 500 nm along the optical axis [10]. This has a major consequence: if several features fall within this region, they will all be flooded with this light at the same time and hence produce signal simultaneously. In the case of fluorescence microscopy, this is the excitation light. As we try to detect the fluorescence signal with a lens and relay it onto a detector, the signals produced by the molecules within this >200-nm spot will be confused. This is because at the detector, each molecule will also produce a spot of focused (fluorescence) light and the spots from these simultaneously illuminated molecules will overlap (Fig. 1.1). No detector will be able to tell the signals from these molecules apart, no matter if it is the eye, a photomultiplier, or even a pixelated camera.

The person who fully appreciated that diffraction poses a serious limit on the resolution was Ernst Abbe, who lived at the end of the nineteenth century and who coined this “diffraction barrier” in an equation which has been named after him [1]. It says that, in order to be separable, two features of the same kind have to be further apart than the wavelength divided by twice the numerical aperture of the objective lens. One can find this equation in most textbooks of physics or optics, and also in textbooks of biochemistry and molecular biology, due to the enormous relevance of light microscopy in these fields. Abbe’s equation is also found on a memorial which was erected in Jena, Germany, where Ernst Abbe lived and worked, and there it is written in stone. This is what scientists believed throughout the twentieth century. However, not only did they believe it, it also was a fact. For example, if one wanted to look at features of the cellular cytoskeleton in the twentieth century [5], this was the type of resolution obtained.

This equation was coined in 1873. So much new physics emerged during the twentieth century and so many new phenomena were discovered. There should be phenomena—at least one—that could be utilized to overcome the diffraction barrier in a light microscope operating with propagating beams of light and regular lenses. S.W.H. understood that it won’t work just by changing the way the light is propagating, the way the light is focused. [Actually he had looked into that; it led him to the invention of the 4Pi microscope [11, 12], which improved the axial resolution, but did not overcome Abbe’s barrier.] S.W.H. was convinced that a potential solution must have something to do with the major discoveries of the twentieth century: quantum mechanics, molecules, molecular states and so on.

Therefore, he started to check his textbooks again in order to find something that could be used to overcome the diffraction barrier in a light-focusing microscope. In simple terms, the idea was to check out the spectroscopic properties of fluorophores, their state transitions, and so on; maybe there is one that can be used for the purpose of making Abbe’s barrier obsolete. Alternatively, there could be a quantum-optical effect whose potential has not been realized, simply because nobody thought about overcoming the diffraction barrier [13].

With these ideas in mind, one day when he was not very far from [Stockholm] in Åbo/Turku, just across the Gulf of Bothnia, on a Saturday morning, S.W.H. browsed a textbook on quantum optics [14] and stumbled across a page that dealt with stimulated emission. All of a sudden he was electrified. Why?

To reiterate, the problem is that the lens focuses the light in space, but not more tightly than 200 nm. All the features within the 200-nm region are simultaneously flooded with excitation light. This cannot be changed, at least not when using conventional optical lenses. But perhaps we can change the fact that all the features which are flooded with (excitation) light are, in the end, capable of sending light (back) to the detector. If we manage to keep some of the molecules dark—to be precise, put them in a non-signaling state in which they are not able to send light to the detector—we will see only the molecules that can, i.e. those in the bright state. Hence, by registering bright-state molecules as opposed to dark-state molecules, we can tell molecules apart. So the idea was to keep a fraction of the molecules residing in the same diffraction area in a dark state, for the period of time in which the molecules residing in this area are detected. In any
case, keep in mind: the state (transition) is the key to making features distinct. And resolution is about discerning features.

For this reason, the question comes up: are there dark states in a fluorescent molecule? The answer is actually contained in the energy diagram in Fig. 1.2b. The ground state of the fluorophore is a dark state! For the molecule to emit fluorescence, the molecule has to be in its excited state. So the excited state is the signaling bright state, but the ground state is, of course, a non-signaling dark state.

What is now the role of stimulated emission? Actually, the answer is as simple as profound: it makes dark molecules, that is, molecules that are not seen by the detector! This was the reason why S.W.H. was so excited. He had found a way to make normal fluorophores not fluoresce, just normal fluorophores that were commonly used in fluorescence microscopy. And now you can easily envisage how the microscope works: stimulated emission depletion—or: STED—microscopy [15–23]. Figure 1.3a sketches the lens, the critical component of a far-field optical

![Diagram](image1.png)

**Fig. 1.2** Switching molecules within the diffraction-limited region transiently “off” (i.e. effectively keeping them in a non-signaling state), enables the separate detection of neighbouring molecules residing within the same diffraction region. (a) In fluorescence microscopy operating with conventional lenses (e.g. confocal microscopy), all molecules within the region covered by the main diffraction maximum of the excitation light are flooded with excitation light simultaneously and emit fluorescence together. This is because they are simultaneously allowed to assume the fluorescent (signalling) state. (b) Keeping most molecules—except the one(s) one aims to register—in a dark state solves the problem. The dark state is a state from which no signal is produced at the detector. Such a transition to the dark “off” state is most simply realized by inducing stimulated emission, which instantaneously forces molecules to their dark (“off”) ground state.
We use a beam of light for exciting molecules from the ground state to the excited state, to make them bright ("on"), i.e. get them to the excited state. Inevitably, the excitation light will be diffracted and one obtains a spot of light of at least 200 nm. Signal which is produced therein, from all the molecules, will be able to end up at the detector. But now, we use a second beam of light which induces stimulated emission, and thus makes dark-state molecules. The idea is to instantly "push" the molecules that were excited back down to the ground state so that the molecule is not capable of emitting light, because it has assumed the dark ground state ("off").

The physical condition for achieving this is that the wavelength of the stimulating beam is longer (Fig. 1.3c). The photons of the stimulating beam have a lower energy, so as not to excite molecules but to stimulate the molecules going from the excited state back down to the ground state. There is another condition, however: we have to ensure that there is indeed a red photon at the molecule which pushes the molecule down. We emphasize this because most red-shifted photons pass by the molecules, as there is a finite interaction probability of the photon with a molecule, i.e. a finite cross-section of interaction. But if one applies a stimulating light intensity at or above a certain threshold, one can be sure that there is at least one photon which "kicks" the molecule down to the ground state, thus making it instantly assume the dark state.
Figure 1.3d shows the probability of the molecule to assume the bright state, the $S_1$, in the presence of the red-shifted beam transferring the molecule to the dark ground state. Beyond a certain threshold intensity, $I_s$, the molecule is clearly turned “off”. One can apply basically any intensity of green light. Yet, the molecule will not be able to occupy the bright state and thus not signal. Now the approach is clear: we simply modify this red beam to have a ring shape in the focal plane [19, 24], such that it does not carry any intensity at the centre. Thus, we can turn off the fluorescence ability of the molecules everywhere but at the centre. The ring or “doughnut” becomes weaker and weaker towards the centre, where it is ideally of zero intensity. There, at the centre, we will not be able to turn the molecules off, because there is no STED light, or it is much too weak.

Now let’s have a look at the sample (Fig. 1.3b) and let us assume that we want to see just the fibre in the middle. Therefore, we have to turn off the fibre to its left and the one to its right. What do we do? We cannot make the ring smaller, as it is also limited by diffraction. Abbe would say: “Making narrower rings of light is not possible due to diffraction.” But we do not have to do that. Rather, we simply have to “shut off” the molecules of the fibres that we do not want to see, that is, we make their molecules dwell in a dark state, until we have recorded the signal from that area. Obviously, the key lies in the preparation of the states. So what do we do? We make the beam strong enough so that the molecules even very close to the centre of the ring are turned “off” because they are effectively confined to the ground state all the time. This is because, even close to the centre of the ring, the intensity is beyond the threshold $I_s$ in absolute terms.

Now we succeed in separation: only in the position of the doughnut centre are the molecules allowed to emit, and we can therefore separate this signal from the signal of the neighbouring fibres. And now we can acquire images with sub-diffraction resolution: we can move the beams across the specimen and separate each fibre from the other, because their molecules are forced to emit at different points in time. We play an “on/off game”. Within the much wider excitation region, only a subset of molecules that are at the centre of the doughnut ring are allowed to emit at any given point in time. All the others around them are effectively kept in the dark ground state. Whenever one makes a check which state they are in, one will nearly always find those molecules in the ground state.

This concept turned out to work very well [17, 19, 23, 25]. Figure 1.4a contains a standard, high-end confocal recording of something which one cannot make out what it is. Figure 1.4b shows the same region imaged using STED microscopy. The resolution is increased by about an order of magnitude (in the red channel), and one can clearly discern what is actually being imaged here: nuclear pore complexes. As a result of the high resolution, it can be seen that this nuclear pore complex features eight molecular subunits. The eightfold symmetry comes out very clearly [25]. There is almost no comparison with the standard confocal recording.

Needless to say, if afforded this increase in spatial resolution, one obtains new information. In other words, new insights are gained with this microscope. Here, we briefly describe research done in collaboration with virologists interested in the human immunodeficiency virus (HIV). Generally, viruses are about 30–150 nm in diameter [5]. So, if one wants to image them with a light microscope … there is no chance this will succeed—one will not see any details of protein distributions on the virus particles. A diffraction-limited fluorescence microscope would yield just a 250–350 nm sized fluorescence blur. The human immunodeficiency virus (HIV) is about 140 nm in size. The collaborating scientists were interested in finding out how a protein called Env is distributed on the HIV particle [26], Fig. 1.5. In the normal recording, nothing specific is seen. In contrast, the high-resolution STED recording revealed that the protein Env forms patterns on the HIV particles. What has actually been found out in this study is that the mature HIV particles—those which are ready to infect the next cell—have the Env concentrated basically in a single place on the virus. It seems to be a requirement for HIV to be very effective—a new mechanistic insight gained as a result of subdiffraction-resolution imaging.
Of course, a strength of light microscopy is that we can image living cells by video-rate recording with STED microscopy. An example are synaptic vesicles in the axon of a living neuron [20]. One can directly see how they move about and we can study their dynamics and their fate over time. It is clearly important to be able to image living cells.

Live-cell imaging “at the extreme” is pictured in Fig. 1.6. Here, we opened the skull of an anaesthetized mouse and looked into the brain of the mouse at the upper, so-called molecular layer of the visual cortex [21]. This was a transgenic mouse, meaning that some of its neurons expressed a fluorescent protein, specifically the yellow fluorescent protein YFP, and this is why this neuron is highlighted from the surrounding brain. The surrounding brain tissue is dark. Next we took sequential recordings and could see the receiving synaptic ends of the neuron—the so-called dendritic spines. They move slightly, and it is worthwhile zooming in on them. One discerns the spine neck and, in particular, the details of the cup-shaped end of the dendritic spines. STED microscopy allows these tiny morphologies to be visualized, such that we can observe their subtle temporal changes. I am very confident that in the not too distant future we will be able to image the proteins here at the synapse [27]. I can also imagine that we will be able to give a visual cue to the

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**Fig. 1.4** Nuclear pore complex architecture in an intact cell nucleus imaged by (a) confocal microscopy (diffraction-limited), and (b) STED nanoscopy. The data was published in [25]
**HIV Envelope protein on single virions**

**Insight:** Env proteins are assembled in mature HIV

**Fig. 1.5** STED nanoscopy of the HIV Envelope protein Env on single virions. Confocal microscopy is not able to reveal the nanoscale spatial distribution of the Env proteins; the images of the Env proteins on the virus particles look like 250–350 nm sized blurred spots (orange, left column). STED microscopy reveals that the Env proteins form spatial patterns (center column, orange), with mature particles having their Env strongly concentrated in space (panel in top row of center column, orange). The data was published in [26]

**Fig. 1.6** STED nanoscopy in living mouse brain. The recording shows a part of a dendrite of a neuron expressing a yellow fluorescent protein (EYFP) in the cytosol, thus highlighting the neuron amidst surrounding (non-labelled) brain tissue. The three to fourfold improved resolution over confocal and multiphoton excitation fluorescence microscopy reveals the dendritic spines (encircled) with superior clarity, particularly the cup-like shape of some of their terminals containing the receiving side of the synapses. The data was published in [21]
mouse and observe how this actually changes the protein distribution directly at the synapse. Thus, in the end we should learn how neuronal communication or memory formation works at the molecular level. Since STED microscopy relies on freely propagating light, one can perform three-dimensional (3D) imaging. It is possible to focus into the brain tissue, for example, and record a 3D data set.

Coming back again to the basics, to the spatial resolution, some will ask: What is the resolution we can get? What is the limit? Indeed, is there a new limit? So let us get back to the principle. The “name of the game” is that we turn off molecules everywhere but at the intensity minimum, at the central zero, of the STED beam [28–31]. If we can make the region in which the molecules are still allowed to emit smaller, the resolution is improved; that is clear. The extent (or diameter) of the region in which the molecules are still “on” now determines the spatial resolution. Clearly, it cannot be described by Abbe’s equation any more. In fact, this diameter must depend on the intensity $I$ which is found at the doughnut crest (Fig. 1.7b, d) and on the threshold intensity $I_s$, which is a characteristic of the photon-molecule interaction. The larger their ratio becomes, the smaller $d$ will become. It is now easy to appreciate that this ratio must be found in the denominator, if we describe the resolution with a new equation which is now obviously required [23, 28, 29]. In fact, $d$ scales inversely with the square root of $I/I_s$. So the larger $I/I_s$, the smaller is $d$ (Fig. 1.8). As a result, $d$ tends to 0 for larger and larger values of $I/I_s$ (Fig. 1.7b, d).

In the situation depicted in Fig. 1.7b, we cannot separate two of the close-by molecules because both are allowed to emit at the same time.
time. But let us make the beam a bit stronger, so that only one molecule “fits in” the region in which the molecules are allowed to be “on”. Now the resolution limit is apparent: it is the size of a molecule, because a molecule is the smallest entity one can separate. After all, we separate features by preparing their molecules in two different states, and so it must be the molecule

Fig. 1.8 Tunable resolution enhancement realized by STED microscopy. (a, b) Confocal (a) and STED (b) image of fluorescent beads with average size of ~24 nm on a cover slip. (c–g) The area of the white rectangle shown in (a) and (b) recorded with different STED intensities. The resolution gain can be directly appreciated. (h) STED depletion η vs. STED-light intensity measured on the same sample. The intensity settings for the measurements (c–g) are marked by red arrows. Scale bars 1 μm (a, b), 200 nm (c–g). Reproduced with permission from [33].

which is the limit of spatial resolution. When two molecules come very close together, we can separate them because at the time one of them is emitting, the other one is “off” and vice versa [28, 30–32].

It is worth noting that if all the “off” or dark molecules are entirely dark, i.e. non-signaling, detecting a single photon from a molecule is absolutely enough to know that there is a molecule present (at the minimum of the STED-beam). The position of that molecule is entirely determined by the presence of the STED-beam photons. These photons determine exactly where the molecule is “on” and where it is “off” (dark). The detected fluorescence photons only indicate the presence of a molecule, or many of them [30–32].

Does one typically obtain molecular spatial resolution, and what about in a cell? For STED microscopy right now, the standard of resolution is between 20 and 40 nm depending on the fluorophore, and depending on the fluorophore’s chemical environment [25]. But this is something which is progressing; it is under continuous development. With fluorophores which have close-to-ideal properties and can be turned “on” and “off” as many times as desired, we can do much better, of course.

In fact, there are such fluorophores—not organic ones, inorganic ones—which meet this requirement already. These are so-called charged nitrogen vacancies in diamond (Fig. 1.9), fluorescent defects in diamond crystals which can be turned on and off an almost unlimited number of times, notably in magnetic sensing and quantum information, which may be eventually read out with diffraction-unlimited spatial resolution using conventional lenses, i.e. even when packed very densely at the nanometer scale.

Fig. 1.9 Fluorophores affording virtually unlimited repetitions of the resolution-enabling on-off state transitions provide the present resolution records in far-field optical imaging using STED, in the single-digit nanometer regime. Color centers (charged nitrogen vacancy centers) in diamond hold great potential for various other applications, notably in magnetic sensing and quantum information.
times [34]. Imaging these, the region of emission was squeezed down to 2.4 nm [35]. It is worth keeping in mind that the wavelength responsible for this result is 775 nm. So the region of emission is smaller than 1%, a very small fraction of the wavelength.

This may look like a proof-of-principle experiment, and to some extent it is. But it is not just that, there is another reason to perform these experiments [34, 36, 37]. The so-called charged nitrogen vacancies are currently regarded as attractive candidates for quantum computation: as qubits operating at room temperature [38, 39]. They possess a spin state with a very long coherence time and which can be prepared and read out optically. Being less than a nanometer in size, they can sense magnetic fields at the nanoscale [40, 41]. There inherently are nanosensors in there, and STED is perhaps the best way of reading out the state and the magnetic fields at the nanoscale. In the end, this could make STED an interesting candidate perhaps for reading out qubits in a quantum computer, or who knows … Development goes on!

Returning to the fundamentals, we have emphasized that the name of the game is “on/off”, or keeping a fraction of the molecules dark for separation [30–32]. This is how we separate molecules, with a bright state and a dark state. Once it is clear that this is a general principle it is obvious that stimulated emission is not the only way by which we can play this “on/off game”. There must also be other “on” and “off” states in a dye which one can use to the same effect [22, 28–30]. With this in mind, S.W.H. browsed other textbooks and found that there are triplet states, long-lived dark states and, of course, in chemistry textbooks, one will find that there is photoinduced cis-trans isomerization (Fig. 1.10). One might ask why use these special transitions that, unlike stimulated emission, are not found in absolutely any fluorophore, as special fluorophores are needed for this? After all, the transitions used in STED are truly basic: optical excitation and de-excitation.

And the two states between which these transitions are induced are the most basic states imaginable, namely the ground and the first excited state.

Indeed, it turns out that there is a strong reason for looking into other types of states and state transitions. Consider the state lifetimes (Fig. 1.10). For the basic STED transition, the lifetime of the state, the excited state, is nanoseconds (Fig. 1.10a). For metastable dark states used in methods termed ground state depletion (GSD) microscopy [42–44] (Fig. 1.10b) the lifetime of the state is microseconds, and for isomerization it is on the order of milliseconds (Fig. 1.10c). Why are these major increases in the utilized state lifetime relevant?

Well, just remember that we separate adjacent features by transferring their fluorescent molecules into two different states. But if the state—one of the states—disappears after a nanosecond, then the difference in states created disappears after a nanosecond. Consequently, one has to hurry up putting in the photons, creating this difference in states, as well as reading it out, before it disappears. But if one has more time—microseconds, milliseconds—one can turn molecules off, read the remaining ones out, turn on, turn off …; they stay there, because their states are long-lived. One does not have to hurry up putting in the light, and this makes this “separation by states” operational at much lower light levels [28, 42].

To be more formal, the aforementioned intensity threshold $I_s$ scales inversely with the lifetime of the states involved (Fig. 1.10e): the longer the lifetime, the smaller is the $I_s$, and the diffraction barrier can be broken using this type of transition at much lower light levels. $I_s$ goes down from megawatts (STED), kilowatts (GSD) down to watts per square centimetre for millisecond switching times—a six orders of magnitude range [28]. This makes transitions between long-lived states very interesting, of course. Here in the equation (Fig. 1.10d), $I_s$ goes down and with that of course also $I$ goes down because one does
not need as many photons per second in order to achieve the same resolution $d$.

The cis-trans isomerization is particularly interesting because it is found in switchable fluorescent proteins. S.W.H. looked into this very early on to check whether it can be used for a STED-like recording. Eventually, S.W.H. called it RESOLFT, for “Reversible Saturable/Switchable Optically Linear (Fluorescence) Transitions” [28, 45–47], simply because he could not have called it STED anymore. There is no stimulated emission in there, which is why he had to give it a different name. The strength is not only that one can obtain high resolution at low light levels. Notably, one can use inexpensive lasers, continuous wave (CW) lasers, and/or spread out the light over a large field of view, because one does not need such intense light to switch the molecules. In this way, one can parallelize the recordings, meaning that one can make an array of many holes (intensity minima, zeros) at the same time and read out a large field of view quickly (Fig. 1.11). It does not matter that one has many of these intensity minima at the same time. As long as they are each further apart than Abbe’s diffraction barrier, they can be read out simultaneously by projecting the signal generated in this array of minima onto a camera. Only a few scanning steps in one direction and in the orthogonal direction, and a super-resolution image of a large field of view is taken. In Fig. 1.12 [48], a living cell was recorded within 2 s with more than 100,000 “doughnuts”, so to speak, in parallel.
Notwithstanding the somewhat different optical arrangement, the key is the molecular transition. Selecting the right molecular transition determines the parameters of imaging. The imaging performance, including the resolution and the contrast level, as well as other factors, is actually determined by the molecular transition chosen [32].

Putting up the next question, what does it take to achieve the best resolution? Now let us assume one had asked this question in the twentieth century. What would have been the answer? Well,
Sure, good lenses. Why? Because the separation of neighbouring features was performed by the focusing of light. Then, of course, one needs good lenses to produce the sharpest focal spot of light at the sample here, there, and everywhere, and/or the sharpest focal spot of light anywhere at the detector. However, once one cannot produce an even smaller focal spot of light, this strategy has come to an end (Fig. 1.13, top). Therefore, if one has several features falling within a diffraction-limited spot of light, one simply cannot do any better. Resolution is definitely limited by diffraction if one separates features by the focusing of light—no way to tell features, the molecules, apart, because everything overlaps on the detector (Fig. 1.13, top). So what was the solution to this problem?

Do not separate just by focusing. Separate by molecular states, in the easiest case by “on/off”-states [28–31]. If separating by molecular states, one can indeed distinguish the features, one can tell the molecules apart even though they reside within the region dictated by diffraction. We can tell, for instance, one molecule apart from its neighbours and discern it (Fig. 1.13, bottom). For
this purpose, we have our choice of states that I have introduced already (Fig. 1.10) which we can use to distinguish features within the diffraction region.

In the methods described, STED, RESOLFT and so on, the position of the state—where the molecule is “on”, where the molecule is “off”—is determined by a pattern of light featuring one or more intensity zeros, for example a doughnut. This light pattern clearly determines where the molecule has to be “on” and where it has to be “off”. The coordinates X, Y, Z are tightly controlled by the incident pattern of light and the position(s) of its zero(s). Moving the pattern to the next position X, Y, Z—one knows the position of the occurrence of the “on” and “off” states already. One does not necessarily require many detected photons from the “on” state molecules, because the detected photons are merely indicators of the presence of a feature. The occurrence of the state and its location is fully determined by the incident light pattern.

Now the question comes up: How does this compare with the seminal invention by Eric Betzig [49], based on the discovery of W. E. Moerner [8, 50], that you can detect single molecules? In the PALM (“Photo-Activated Localization Microscopy”) [49] concept (also called STORM or FPALM [51, 52]), there are two fundamental differences to STED-like approaches (Fig. 1.14). First of all, it critically relies on the detection of single molecules. Secondly, unlike in the STED case, in the PALM case the spatial position of the on-state is uncontrollable, totally stochastic. A molecule “pops up” somewhere randomly in space, a single molecule per diffraction-sized region, and it is in this way that the “on”/“off” state difference is created. But since one does not know where a molecule has turned to the on-state, a pattern of light must be...
used with which one can measure the position. This pattern of light is the fluorescent light which is emitted by the molecule and imaged onto an array-detector, usually a camera. The pixels of the camera provide the coordinate reference. Without going into the details, this pattern of emitted fluorescence light allows one to determine the molecule’s position with a centroid calculation.

An interesting insight here is that one needs a bright pattern of emitted light to find out the position just as one needs a bright pattern of incident light in STED/RESOLFT to determine the position of emission. Not surprisingly, one *always* needs bright patterns of light when it comes to positions, because if one has just a single photon, this alone tells nothing. The photon can go anywhere within the realm of diffraction, there is no way to control where it goes within the diffraction zone. In other words, when dealing with positions, one needs *many* photons by definition, because this is inherent to diffraction. Many photons are required for defining positions of “on”- and “off”-state-molecules in STED/RESOLFT microscopy, just as many photons are required to find out the position of “on”-state molecules in the stochastic method PALM.

One is not confined to using a single doughnut (a single diffraction zone) in STED/RESOLFT. We can use a “widefield” arrangement, meaning that we can also record a large field of view (compare the blue pattern in Fig. 1.11). To this end, we parallelize the scanning using an array of intensity minima, such as an array of doughnuts. Again, the fundamental difference to the spatially stochastic methods is (Fig. 1.15) that the positions where the molecules can assume the “on”- or the “off”-state are tightly controlled by the pattern of light with which we illuminate the sample. This is regardless of whether there is one molecule at the intensity minimum of the pattern, or three molecules; however many, it does not matter.

Although the PALM principle can also be implemented on a single diffraction zone only (i.e. using a single focused beam of light), it is usually implemented in a “parallelized” way, i.e. on a larger field of view containing many diffraction zones. PALM parallelization requires that there may be only a single “on”-state molecule within a diffraction zone, i.e. within the distance dictated by the diffraction barrier. However, the position of this molecule is completely random. Therefore, we have to make sure that the “on”-state molecules are far enough apart from each other, so that they are still identifiable as separate molecules. While in (STED/RESOLFT) the position of a certain state is given by the pattern of light falling on the sample, position in PALM is established from the pattern of (fluorescence) light coming out of the sample.

What does $I/I_s$ in STED/RESOLFT stand for? $I_s$ can be seen as the number of photons that one needs to ensure that there is at least one photon interacting with the molecule, pushing it from one state to the other in order to create the required difference in molecular states. $I/I_s$ is, so to speak, the number of photons which really “can do something” at the molecule while most of the others just “pass by”. Similarly, in the PALM concept, the number of photons $n$ in $1/\sqrt{n}$ is the number of those photons that are detected, i.e. that really contribute to revealing the position of the emitting molecule. In other words, in both concepts, to attain a high coordinate precision, one needs *many* photons that really do something. This analogy very clearly shows the importance of the number photons to achieve coordinate precision in both concepts.

However, in both cases the separation of features is, of course, accomplished by an “on/off”-transition [28–31]. This is how we make features distinct, how we tell them apart. As a matter of fact, all the super-resolution methods which are in place right now and really useful, achieve molecular distinguishability by transiently placing the molecules that are closer together than the diffraction barrier in two different states for the time period in which they are jointly scrutinized by the detector. “Fluorescent” and “non-fluorescent” is the easiest pair of states to play with, and so this is what has worked out so far.

One can take the point of view that in the twentieth century it was the lenses which were decisive. And the lens makers ruled the field. One had to go to them and ask them for...
the best lenses to get the best resolution. But how is it today? No, it is not the lens makers. This resolution game is not about lenses anymore. It is about molecular states, and molecular states are of course about molecules. The molecules determine now how well we can image; they determine the spatial resolution. And that is not optical technology—that is chemistry. In a way this was initially a physics problem—the diffraction barrier certainly was, no doubt about it—which has now evolved into a chemistry topic.

The enabling element being a transition between two states, the two states need not be fluorescence “on”/“off”: they could also be a pair of states “A” and “B”, like “absorption/non-absorption”, “scattering/non-scattering”, “spin up/spin down”, “bound/unbound” (as in the method called PAINT [53]), etc. Perhaps one can also imagine a super-resolution absorption microscope or a super-resolution scattering microscope, if one identifies the right states.

The field is progressing rapidly, and some selected highlights have been the demonstration of STED nanoscopy in 3D [54, 55] (compare Fig. 1.16),

**Fig. 1.15** To parallelize STED/RESOLFT scanning, a “widefield” arrangement with an array of intensity minima (e.g. an array of doughnuts) may be used. The numbers of molecules at these readout target coordinates do not matter, while PALM requires that there may be only a single “on”-state molecule within a diffraction zone, i.e. within the distance dictated by the diffraction barrier. [More precisely: the number of molecules per diffraction zone has to be so low that each molecule is recognized individually.] The position of each on-state molecule is however completely random in space. \( I \), can be regarded as the number of photons that one needs to ensure that there is at least one photon interacting with the molecule, pushing it from one state to the other in order to create the required difference in molecular states. \( III \), is, so to speak, the number of photons which really elicit the (on/off) state transition at the molecule, while most of the others just “pass by”. Similarly, in the PALM concept, the number of photons \( n \) in \( 1/\sqrt{n} \) is the number of those photons that are really detected at the coordinate-giving pixelated detector (camera), i.e. that really contribute to revealing the position of the emitting molecule. In other words, in both concepts, to attain a high coordinate precision, one needs many photons that act. The references shown are to [8, 9, 49, 51, 52] described in the text.
Fig. 1.16 3D STED microscopy for simultaneously increasing the resolution in the focal plane and along the optic axis. **(Left Top)** Schematic setup. The STED power is distributed between the two phase plates (P_{lat} and P_{3D}) by using a combination of a λ/2 plate and a polarizing beam splitter (PBS). The second PBS recombines the two beams incoherently. The excitation (Exc) and STED beams are overlaid by a dichroic mirror (DM). A λ/4 plate ensures the circular polarization of all beams prior to being focused by the objective lens (OL). The fluorescence signal (Fl) is collected by the same lens. **(Left Bottom)** Focal intensity distributions of excitation and STED beams measured using gold beads in reflectance mode. From left to right: Excitation, STED beam from P_{lat} arm resulting in the focal deexcitation pattern STED_{lat}, STED beam from P_{3D} arm yielding STED_{3D}, incoherent combination of both arms (30% STED_{lat}/70% STED_{3D} power distribution). The latter distribution results in an efficient coverage of the volume around the focal point. Scale bars 500 nm. **(Right)** (a–d) 3D nanoscale image of dilute distribution of 20 nm diameter fluorescent spheres on glass. xy sections of (a) confocal and (b) STED. (c) Confocal and (d) STED xz sections along the dashed blue line indicated in panels (a) and (b). Individual beads can be easily resolved in the STED images. Comparing panel (c) with panel (d), note the significant reduction in cross-sectional area in the STED xz-image (e, f) Intensity profile along the (e) x and (f) z direction for sections indicated by the white arrows in panels (c) and (d). All presented data is raw data. (g) Focal volume reduction relative to confocal focal volume measured using 20 nm fluorescent spheres. The combination of two de-excitation patterns gives a maximal volume reduction factor of 125. Scale bars 1 μm. From [54], reproduced with permission.
Fig. 1.17 isoSTED: Fluorescence microscopy setup with isotropic 3D focal spot. (a) Beams for excitation, STEDxy (lateral) and STEDz (axial) are combined using a dichroic mirror (DCSP) and then fed through a beam scanner into a 4Pi unit with two opposing objective lenses (O1 and O2; HCX PL APO 100x, 1.46 NA OIL CORR). The fluorescence light (orange) collected by both lenses backpropagates along the same optical path to the detector, passing through the DCSP and a second dichroic mirror (DCLP). The pivot plane (PP) of all scanning beams is conjugated to the entrance pupils of the objective lenses. The incoming beams are divided by a polarizing beam-splitter (PBS) and coherently superimposed at both lenses’ common focal plane inside the sample (S). A piezo-driven mirror (MP) controls the difference in pathlength between both cavity arms and thereby the 4Pi phases of all beams. The polarization state of STEDxy and STEDz is adjusted by two half-wave retarder plates (H1 and H2). The excitation beam and the STED beams for lateral (STEDxy, imprinted with a circular phase ramp (PM)) and axial (STEDz) fluorescent spot compression are polarized under $\alpha = 45^\circ$ with respect to the perpendicular direction (n) to the splitting plane (p) of the polarizing beam-splitter. STEDxy and STEDz are polarized orthogonal to each other. (b) Calculated focal intensity distributions and formation of the STED PSF with respective wavelengths, $\lambda$, and 4Pi phases $\phi$. (c) Isotropic effective focal spot (PSF) on the nanoscale. (Left) Calculated PSF of a confocal fluorescence microscope and the corresponding spherical PSF of the isoSTED microscope at $L_0/L_1 = 15$ (NA = 1.4). (Middle) Experimental counterpart to (left) as measured with a 21-nm-diameter fluorescence bead. The FWHM of the confocal setup (1.46 NA) is 165 nm in the lateral and 405 nm in the axial direction. Switching on the STED pattern shown in (b) leads to a largely spherical main focal fluorescence spot. (Right) Gaussian fits through the lateral and axial profiles of the focal spot yield indicated FWHM, corresponding to an isotropic far-field optical 3D resolution of $\lambda/16$. Baselines are marked with colored circles. Scale bars, 250 nm. From [55], reproduced with permission.
Fig. 1.18 Lightsheet (LS)-RESOLFT concept. A living specimen expressing RSFPs is grown on a coverslip mounted on a movable platform. The specimen is illuminated (here in y direction) perpendicular to the detection axis (z). Only in a thin diffraction-limited section, RSFPs are switched from their initial off state (unfilled dots) to the on state (white dots) by an activating LS. None of the fluorophores outside the illuminated volume is affected by the laser light. An LS featuring a central zero-intensity plane switches off the activated RSFPs above and below the detection focal plane (x–y). For negative-switching RSFPs, this is a competing process to fluorescence (green dots and arrows). For off-switching light intensities above the threshold of the RSFPs, only fluorophores within a slice of subdiffraction thickness remain activated. These can be read out by a third LS and contribute to the LS-RESOLFT image. The platform is displaced to the next position in the scanning sequence for another illumination cycle. The lower row shows measured y–z cross-sections of the applied LSs visualized in fluorescent medium. The sheets impinge on the coverslip at an angle of 30°. (Scale bar, 100 μm.) From [57], reproduced with permission.

at millisecond imaging times for ultrafast dynamics in small fields of view [56], the demonstration of a RESOLFT strategy to neutralize the diffraction limit of light-sheet fluorescence microscopy [57] (compare Fig. 1.18), efficient STED nanoscopy with quantum dots as fluorescent labels [58]; highest levels of 3D isotropic resolution (<30 nm in x, y, z simultaneously) with a new, stable design of 4Pi-based isoSTED [59] (compare Fig. 1.17 for more details on the isoSTED approach), and the several-thousand fold “massive” parallelization of RESOLFT and even STED without resolution compromises for faster imaging of large fields [60, 61] as well as extended multicolor capabilities [62] and nanoscopy in living animals (Fig. 1.19).
**Fig. 1.19** Super-resolution microscopy *in vivo*: mouse and fruit fly nanoscopy. (a) STED nanoscopy of a mouse with enhanced yellow fluorescent protein-labelled neurons. Shown are dendritic and axonal details in the molecular layer of the somatosensory cortex of a living, anesthetized mouse. Optical access to the brain cortex was enabled by a cover glass-sealed cranial window. Top panel: image of a neuron. Bottom panel: STED time-lapse recording of spine morphology dynamics. Scale bars: 1 μm. (b) STED imaging of synaptic protein distribution. Example: PSD95, the abundant scaffold protein at the postsynaptic membrane, which organizes numerous other synaptic proteins. (Left) The cartoons show the *in vivo* labeling of endogenous PSD95-HaloTag, a self-labeling enzymatic protein tag, with organic fluorophores. (Right) Depending on the orientation of the individual spine head imaged with respect to the focal plane, the intricate spatial organization of PSD95 at the synapse is revealed in the STED mode. Scale bars: 500 nm. (c) RESOLFT imaging of the microtubule cytoskeleton of intact, living Drosophila melanogaster larvae. A second instar larva ubiquitously expressing a fusion protein composed of the reversibly switchable fluorescent protein (RSFP) rsEGFP2 fused to α-tubulin was placed under a coverslip and imaged through the intact cuticle. Left: confocal overview. Middle and right: magnifications of the area indicated by the corresponding square. Shown are comparisons of confocal and RESOLFT recordings (separated by a dashed line), exemplifying the difference in resolution. Scale bars: 10 μm, 1 μm and 500 nm (from left to right). Part (a) is adapted from [21]. Reprinted with permission from AAAS. Part (c) is adapted with permission from [63], CC-BY 3.0. Parts (a) and (c) reproduced with permission from [64], part (b) from [65].
1.2 Recent Developments: Nanoscopy at the MINimum

Improvements to STED microscopy have substantially expanded its capabilities for a growing diversity of cell-biological applications [64]. Concretely, recent adaptive scanning strategies [66–68] have proven key to reducing the overall light dose applied to the sample. These conceptual additions to STED/RESOLFT imaging reduce photobleaching [69] and are advantageous for live-cell imaging. Thus, they have allowed the resolution of STED microscopy to be pushed even closer to the <20 nm regime for organic fluorophores and for routine users, under realistic cell-imaging conditions. The first example of these approaches is MINFIELD [67], which provides major signal increases and prolonged acquisitions by restricting imaging to regions below the diffraction limit (Fig. 1.20a). MINFIELD STED microscopy avoids the exposure of the fluorophore to excess intensities of the ‘doughnut’ and, more generally, to the maxima of the light intensity distribution used for on/off-switching. Rapid and repetitive MINFIELD recording is likely to be the approach of choice for investigating small spatial domains, such as the synapse. Moreover, MINFIELD STED microscopy will allow fast dynamical nanoscale processes to be captured on millisecond timescales and beyond [56, 67].

DyMIN is a related recording strategy [68] which minimizes exposure to unduly high intensities except at scanning steps where these intensities are strictly required for resolving features (Fig. 1.20b–d). Like MINFIELD, the DyMIN approach achieves dose reductions by up to orders of magnitude, particularly for relatively sparse fluorophore distributions. Initially demonstrated for STED immunofluorescence imaging, both MINFIELD and DyMIN will be explored for other classes of fluorophores, including the inherently lower-light-level RESOLFT nanoscopy variants with genetically encoded fluorescent proteins. The recently described organic switchable photochromic compounds [70] will also be further developed as attractive alternatives in this regard. The synergistic combination of two separate fluorophore state transitions in a recent concept termed multiple off-state transitions (MOST) for nanoscopy [66] has also enabled many more image frames to be captured, at much improved contrasts and with lower STED light dose at a given resolution than for standard STED. Approaches to directly count molecules with STED have also been developed [71], and can be used to quantify the composition of suitably labeled molecular clusters.

Of all the nanoscopy or super-resolution advances of the last decade, the recently described MINFLUX concept [72] stands out, because it contains a radically new idea. Whereas in PALM/STORM the localization of a molecule is based on maximizing the number of detected fluorescence photons on a camera, which is inevitably limited by bleaching, in MINFLUX (Figs. 1.21 and 1.22) the molecule is localized by making it coincide with the intensity zero of a doughnut-shaped excitation beam. The excitation beam is scanned across the molecule and the fluorescence is typically recorded as in a confocal microscope. The position of the molecule is ultimately identical with the position of the doughnut at which fluorescence emission is minimal (see Fig. 1.21). By fundamentally reducing the number of detected photons required for nanometer-precise localization, MINFLUX has opened the door to low-light level optical analysis of tiny objects at true molecular scale resolution (1–5 nm). With MINFLUX, lens-based fluorescence microscopy has thus reached the ultimate resolution limit: the size of the fluorescent molecule itself. Moreover, the resolution is attained at relatively high speed, at least 10 times faster than in PALM/STORM.

While the experimental developments of the MINFLUX concept are still in the beginnings, it is worth commenting on the fundamental advantage over localization based on the emitted fluorescence alone. As discussed in [72, 73], in PALM/STORM, as in camera-based tracking applications, a molecule’s position is inferred from the maximum of its fluorescence diffraction pattern (back-projected into sample space). The precision of such camera-based localization ideally reaches $\sigma_{\text{cam}} \geq \sigma_{\text{PSF}}/\sqrt{N}$, with $\sigma_{\text{PSF}}$ being the standard deviation of the pattern and $N$ the num-
Fig. 1.20 Concepts with improved sample-responsive implementation of the on-off switching. (a) The MINFIELD concept: Lower local de-excitation intensities in STED nanoscopy for image sizes below the diffraction limit. (Left) In STED imaging with pulsed lasers, the ability of a fluorophore to emit fluorescence decreases nearly exponentially with the intensity of the beam de-exciting the fluorophore by stimulated emission, $I$, can be defined as the intensity at which the fluorescence signal is reduced by 50%. Fluorophores delivering higher signal are defined as on, whereas those with smaller signal are defined as off. (Middle) The STED beam is shaped to exhibit a central intensity zero in the focal region (i.e., a doughnut), so that (Right) molecules can show fluorescence only if they are located in a small area in the doughnut center. This area decreases with increasing total doughnut intensity. Due to its diffraction-limited nature, the intensity distribution of the STED focal beam extends over more than half of the STED-beam wavelength and exhibits strong intensity maxima, significantly contributing to photobleaching. By reducing the size of the image field to an area below the diffraction limit, where the STED beam intensity is more moderate (i.e., around the doughnut minimum; compare image area indicated in the Middle), one can reduce the irradiation intensities in the area of interest, inducing lower photobleaching and allowing the acquisition of more fluorescence signal at higher resolution. Scale bar: 200 nm. (b) DyMIN (Dynamic Intensity MINimum) STED imaging. (Left) Concept illustrated for two fluorophores spaced less than the diffraction limit. Signal is probed at each position, starting with a diffraction-limited probing step ($P_{\text{STED}} = 0$, Top), followed by probing at higher resolution ($P_{\text{STED}} > 0$). At any step, if no signal indicates the presence of a fluorophore, the scan advances to the next position without applying more STED light to probe at higher resolution. For signal above a threshold (e.g., $T_1$, Upper Middle), the resolution is increased in steps (Lower Middle), with decisions taken based on the presence of signal. This is continued up to a final step of $P_{\text{max}}$ (full resolution where required). For the highest-resolution steps, directly at the fluorophore(s), the probed region itself is located at the minimum of the STED intensity profile (Bottom). (c) Dual-color isotropic nanoscopy of nuclear pore components and lamina with DyMIN STED: Confocal and 3D DyMIN STED recordings of nuclear pore complexes (shown in green) and lamina (red). Scale bars: 500 nm. (d) DyMIN STED imaging of DNA origami structures with fluorophore assemblies. The DNA origami-based nanorulers with nominally 30-nm separation (10-nm gap) consisted of two groups of ~15 ATTO647N fluorophores, on average, each. Accounting for the known ~20-nm extent of the fluorophore groups (compare schematic), the widths of the Gaussians imply an effective PSF of ~17 nm (FWHM). Scale bars: 200 nm. Figures reproduced with permission from [67] (a) and [68] (b–d)
Fig. 1.21 Principles of MINFLUX, a concept for localizing photon emitters in space, illustrated in a single dimension (x) by using a standing light wave of wavelength \( \lambda \). (a) The unknown position \( x_m \) of a fluorescent molecule is determined by translating the standing wave, such that one of its intensity zeros travels from \( x = -L/2 \) to \( L/2 \), with \( x_m \) somewhere in between. (b) Because the molecular fluorescence \( f(x) \) becomes zero at \( x_m \), solving \( f(x_m) = 0 \) yields the molecular position \( x_m \). Equivalently, the emitter can also be located by exposing the molecules to only two intensity values belonging to functions \( I_0(x) \) and \( I_1(x) \) that are fixed in space having zeros at \( x = -L/2 \) and \( L/2 \), respectively. Establishing the emitter position can be performed in parallel with another zero, by targeting molecules further away than \( \lambda/2 \) from the first one. (c) Localization considering the statistics of fluorescence photon detection: Success probability \( p_0(x) \) for various beam separations \( L \) are shown as listed in the legend for \( \lambda = 640 \) nm. The fluorescence photon detection distribution \( P(n_0|N = n_0 + n_1 = 100) \) conditioned to a total of 100 photons is plotted along the right vertical axis of normalized detections \( n_0/N \) for each \( L \). The distribution of detections is mapped into the position axis \( x \) through the corresponding \( p_0(x,L) \) function (gray arrows), delivering the localization distribution \( P(x_m|N = 100) \). The position estimator distribution contracts as the distance \( L \) is reduced. (d) Cramér-Rao bound (CRB) for each \( L \). Precision is maximal halfway between the two points where the zeros are placed. For \( L = 50 \) nm, detecting just 100 photons yields a precision of 1.3 nm. Figure reproduced from [72]. Reprinted with permission from AAAS.
**Fig. 1.22** The MINFLUX concept: molecular resolution in fluorescence nanoscopy. (a) Implementation of MINFLUX in 2D fluorescence imaging and tracking. (Top) Diagrams of the positions of the doughnut in the focal plane and resulting fluorescence photon counts. (Bottom) Basic application modalities of MINFLUX. (Left) Nanoscopy: A nanoscale object features molecules whose fluorescence can be switched on and off, such that only one of the molecules is on within the detection range. They are distinguished by abrupt changes in the ratios between the different n_0,1,2,3 or by intermissions in emission. (Middle) Nanometer-scale (short-range) tracking: The same procedure can be applied to a single emitter that moves within the localization region of size L. As the emitter moves, different fluorescence ratios are observed that allow the localization. (Right) Micron-scale (long-range) tracking: If the emitter leaves the initial L-sized field of view, the triangular set of positions of the doughnut zeros is (iteratively) displaced to the last estimated position of the molecule. By keeping it around r_0 by means of a feedback loop, photon emission is expected to be minimal for r_0 and balanced between n_1, n_2, and n_3, as shown. (b) With MINFLUX nanoscopy one can, for the first time, separate molecules optically which are only a few nanometers apart from each other. On the left, a schematic of the molecules is presented. Whereas the ultra-high resolution PALM/STORM microscopy at the same molecular brightness (Right) delivers a diffuse image of the molecules (here in a simulation under ideal technical conditions), the position of the individual molecules can be easily discerned with the practically realized MINFLUX (middle). (c) Many much faster movements can be followed than is possible with STED or PALM/STORM microscopy. Left: Movement pattern of 30S ribosomes (colored) in an *E. coli* bacterium (gray scale). Right: Movement pattern of a single 30S ribosome (green) shown enlarged. (d) MINFLUX tracking of rapid movements of a custom-designed DNA origami. (Top left) Diagram of the DNA origami construct with a single ATTO 647N fluorophore attached at the center of the bridge (10 nm from the origami base). By design, the emitter can move on a half-circle above the origami and is thus ideally restricted to a 1D movement. (Bottom left) Histogram of 6118 localizations of the sample with δt = 400 μs time resolution and a 1.5 × 1.5-nm binning. The predominant motion is along a single direction. (Right, Upper) A 300-ms excerpt of the photon count trace (time resolution δt = 400 μs per localization). The color coding corresponds to the zero positions shown to the left. (Right, Lower) Mean-subtracted trajectory. Figure reproduced from [72] (a) (reprinted with permission from AAAS) and [73] (d)
ber of fluorescence photons making up the pattern [74]. Note that $\sigma_{\text{cam}}$ is thus clearly bounded by the finite fluorescence emission rate, which for currently used fluorophores rarely allows more than a few hundred photon detections per millisecond ($<1$ MHz). Moreover, emission is frequently interrupted and eventually ceases due to blinking and bleaching. This also keeps the photon emission rate as the limiting factor for the obtainable spatio-temporal resolution. As a result, state-of-the-art single-molecule tracking performance long remained in the tens of nanometer per several tens of millisecond range.

Drawing on the basic ideas of the coordinate determination employed in STED/RESOLFT microscopy, the MINFLUX concept addresses these fundamental limitations [72]. By localizing individual emitters with an excitation beam featuring an intensity minimum that is spatially precisely controlled, MINFLUX takes advantage of coordinate targeting for single-molecule localization. The basic steps are illustrated for one spatial dimension in Fig. 1.21. In a typical two-dimensional MINFLUX implementation, the position of a molecule is obtained by placing the minimum of a doughnut-shaped excitation beam at a known set of spatial coordinates in the molecule’s proximity. These coordinates are within a range $L$ in which the molecule is anticipated (Fig. 1.22a). Probing the number of detected photons for each doughnut minimum coordinate yields the molecular position. It is the position at which the doughnut would produce minimal emission, if the excitation intensity minimum were targeted to it directly. As the intensity minimum is ideally a zero, it is the point at which emission is ideally absent. The precision of the position estimate increases with the square root of the total number of detected photons and, more importantly, by decreasing the range $L$, the spatial scale inserted from the outside into the experiment. For small ranges $L$ for which the intensity minimum is approximated by a quadratic function, the localization precision does not depend on any wavelength and, for the case of no background and perfect doughnut control, the precision $\sigma_{\text{MINFLUX}}$ simply scales with $L/\sqrt{N}$ at the center of the investigated range. In other words, the better the coordinates of the excitation minimum match the position of the molecule, the fewer fluorescence detections are needed to reach a given precision.

In the conceptual limit where the excitation minimum coincides with the position of the emitter, i.e. $L = 0$, the emitter position is rendered by vanishing fluorescence detection. This is contrary to conventional centroid-based localization where precision improvements are tightly bound to having increasingly larger numbers of detected photons.

The already demonstrated tracking of fluorophores with substantially sub-millisecond position sampling (Fig. 1.22c) is only the beginning in a quest for highest spatiotemporal capabilities (compare data in Fig. 1.22d) [73]. The inherent confocality should also provide a critical advantage when considering imaging in more dense and three-dimensional specimens, such as brain slices and in-vivo imaging scenarios. With further development of other aspects, including field-of-view enlargement, etc., MINFLUX is bound to transform the limits of what can be observed in cells and molecular assemblies with light. This should most probably impact cell and neurobiology and possibly also structural biology. Moreover, it should be a great tool for studying molecular interactions and intra-macromolecular dynamics in a range that has not been accessible so far.

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References

1. Abbe E. Beiträge zur Theorie des Mikroskops und der mikroskopischen Wahrnehmung. Arch Mikros Anat. 1873;9:413–68.
2. Verdet E. Leçons d’optique physique, vol. 1. Paris: Victor Masson et fils; 1869.
3. Lord Rayleigh. On the theory of optical images, with special reference to the microscope. Philos Mag. 1896;5(42):167–95.
4. von Helmholtz H. Die theoretische Grenze für die Leistungsfähigkeit der Mikroskope. Ann Phys Chem. 1874;6:557–84.
5. Alberts B, et al. Molecular biology of the cell. 4th ed. New York, NY: Garland Science; 2002.
6. Giepmans BNG, et al. The fluorescent toolbox for assessing protein location and function. Science. 2006;312(5771):217–24.
7. Lakowicz JR. Principles of fluorescence spectroscopy. New York, NY: Springer; 2006.
8. Moerner WE, Kador L. Optical-detection and spectroscopy of single molecules in a solid. Phys Rev Lett. 1989;62(21):2535–8.
9. Orrit M, Bernard J. Single pentacene molecules detected by fluorescence excitation in a p-terphenyl crystal. Phys Rev Lett. 1990;65:2716–9.
10. Born M, Wolf E. Principles of optics. 7th ed. Cambridge; New York, NY, Melbourne, VIC; Madrid, Cape Town: Cambridge University Press; 2002.
11. Hell SW, Stelzer EHK. Properties of a 4pi confocal fluorescence microscope. J Opt Soc Am. 1992;9:2159–66.
12. Hell SW, Schrader M, Van der Voort HTM. Far-field fluorescence microscopy with three-dimensional resolution in the 100-nm range. J Microsc. 1997;187(1):1–7.
13. Hell SW. Improvement of lateral resolution in far-field light microscopy using two-photon excitation with offset beams. Opt Commun. 1994;106:19–24.
14. Loudon R. The quantum theory of light. Oxford: Oxford University Press; 1983.
15. Hell SW, Wichmann J. Breaking the diffraction resolution limit by stimulated-emission - stimulated-emission-depletion fluorescence microscopy. Opt Lett. 1994;19(11):780–2.
16. Klar TA, Hell SW. Subdiffraction resolution in far-field fluorescence microscopy. Opt Lett. 1999;24(14):954–6.
17. Klar TA, et al. Fluorescence microscopy with diffraction resolution barrier broken by stimulated emission. Proc Natl Acad Sci U S A. 2000;97:8206–10.
18. Donnert G, et al. Macromolecular-scale resolution in biological fluorescence microscopy. Proc Natl Acad Sci U S A. 2006;103(31):11440–5.
19. Willig KI, et al. STED microscopy reveals that synaptotagmin remains clustered after synaptic vesicle exocytosis. Nature. 2006;440(7086):935–9.
20. Westphal V, et al. Video-rate far-field optical nanoscopy dissects synaptic vesicle movement. Science. 2008;320(5873):246–9.
21. Berning S, et al. Nanoscopy in a living mouse. Science. 2012;335(6068):551.
22. Dyba M, Hell SW. Focal spots of size lambda/23 open up far-field fluorescence microscopy at 33 nm axial resolution. Phys Rev Lett. 2002;88(16):163901.
23. Westphal V, Hell SW. Nanoscale resolution in the focal plane of an optical microscope. Phys Rev Lett. 2005;94:143903.
24. Keller J, Schoenle A, Hell SW. Efficient fluorescence inhibition patterns for RESOLFT microscopy. Opt Express. 2007;15(6):3361–71.
25. Göttfert F, et al. Coaligned dual-channel STED nanoscopy and molecular diffusion analysis at 20 nm resolution. Biophys J. 2013;105(1):L01–3.
using reversibly photoswitchable proteins. Proc Natl Acad Sci U S A. 2005;102(49):17565–9.
46. Grotjohann T, et al. Diffraction-unlimited all-optical imaging and writing with a photochromic GFP. Nature. 2011;478(7368):204–8.
47. Hell SW, Dyba M, Jakobs S. Concepts for nanoscale resolution in fluorescence microscopy. Curr Opin Neurobiol. 2004;14(5):599–609.
48. Chmyrov A, et al. Nanoscopy with more than 100,000 ‘doughnuts’. Nat Methods. 2013;10(8):737–40.
49. Betzig E, et al. Imaging intracellular fluorescent proteins at nanometer resolution. Science. 2006;313(5793):1642–5.
50. Dickson RM, Cubitt AB, Tsien RY, Moerner WE. Ultra-high resolution imaging by fluorescence photoactivation localization microscopy. Biophys J. 2006;91(11):4258–72.
51. Sharonov A, Hochstrasser RM. Wide-field subdiffraction imaging by accumulated binding of diffusing probes. Proc Natl Acad Sci U S A. 2006;103:18911–16.
52. Harke B, Ullal CK, Keller J, Hell SW. Three-dimensional nanoscopy of colloidal crystals. Nano Lett. 2008;8(5):1309–13.
53. Schmidt R, Wurm CA, Jakobs S, Engelhardt J, Egner A, Hell SW. Spherical nanosized focal spot unravels the interior of cells. Nat Methods. 2008;5(6):539–44.
54. Schneider J, Zahn J, Maglione M, Sigrist SJ, Marquard J, Chojnacki J, Kräusslich H-G, Sahl SJ, Engelhardt J, Hell SW. Ultrafast, temporally stochastic STED nanoscopy of millisecond dynamics. Nat Methods. 2015;12(9):827–30.
55. Hoyer P, de Medeiros G, Balázs B, Norlin N, Besir C, Hanne J, Kräusslich H-G, Sahl SJ, Engelhardt J, Hell SW. STED fluorescence microscopy by imaging regions of subdiffraction extent. Proc Natl Acad Sci U S A. 2016;113(13):3442–6.
56. Hanne J, Falk HJ, Görlich F, Hoyer P, Engelhardt J, Sahl SJ, Hell SW. STED nanoscopy with fluorescent quantum dots. Nat Commun. 2015;6:4273.
57. Curdt F, Herr SJ, Lutz T, Schmidt R, Engelhardt J, Sahl SJ, Hell SW. isoSTED nanoscopy with intrinsic beam alignment. Opt Express. 2015;23(4):30891–903.
58. Berghauss F, Alber L, Sahl SJ, Engelhardt J, Hell SW. 2000-fold parallelized dual-color STED fluorescence nanoscopy. Opt Express. 2015;23(1):211–23.
59. Chmyrov A, Leutenegger M, Grotjohann T, Schönhle A, Keller-Findeisen J, Kastrup L, Jakobs S, Donnert G, Sahl SJ, Hell SW. Achromatic light patterning and improved image reconstruction for parallelized RESOLFT nanoscopy. Sci Rep. 2017;7:44619.
60. Winter FR, Loidolt M, Westphal V, Butkevich AN, Gregor C, Sahl SJ, Hell SW. Multicolour nanoscopy of fixed and living cells with a single STED beam and hyperspectral detection. Sci Rep. 2017;7:46492.
61. Thompson RE, Larson DR, Webb WW. Precise nanometer localization analysis for individual fluorescent probes. Biophys J. 2002;82(5):2775–83.
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