Review Article

Benefits from Treatment and Control of Patients with Resistant Hypertension

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Resistant hypertension is commonly found in everyday clinical practice. However, the risks of resistant hypertension, as well as the benefits of treatment and control of blood pressure in patients with resistant hypertension remain vaguely clarified. Data from small clinical studies and observational cohorts suggest that patients with resistant hypertension are at increased cardiovascular risk, while control of blood pressure offers substantial benefits. It has to be noted however that data from appropriate large randomized studies are missing, and resistant hypertension remains remarkably understudied. Resistant hypertension has attracted significant scientific interest lately, as new therapeutic modalities become available. The interventional management of resistant hypertension either by carotid baroreceptor stimulation or renal sympathetic denervation is currently under investigation with promising preliminary results. This review presents available evidence regarding the benefits of treatment and control of blood pressure in patients with resistant hypertension and offers a critical evaluation of existing data in this field.

1. Introduction

Resistant hypertension is defined as uncontrolled blood pressure despite the use of optimal doses of three antihypertensive medications, of which one is a diuretic [1]. Although this definition encompasses a large number of patients, many of these patients can be controlled with more careful adjustment of their regimen and implementation of good practices. Several factors have been identified as contributors to resistant hypertension: poor patient adherence, physician inertia, inadequate doses or inappropriate combinations of antihypertensive drugs, secondary forms of hypertension, drug-induced hypertension, excess alcohol intake, and volume overload [2]. Lifestyle modifications including salt restriction are very important in these patients [3]. Addressing some of the comorbid conditions, such as sleep apnea, primary aldosteronism [4], or addition of adjunct therapies such as spironolactone [5–11] can achieve blood pressure control. However, many patients remain uncontrolled despite the use of four, five, or six antihypertensive drugs, especially in everyday clinical practice, outside the “sterile” environment of clinical trials. It is surprising to realize that although hypertension is among the most studied diseases, resistant hypertension which denotes the most severe, high-risk, and probably more scientifically interesting subgroup remains so much understudied.

Unfortunately, data regarding the natural history of resistant hypertension is limited. Furthermore, the benefits of controlling blood pressure in patients with resistant hypertension are vaguely clarified, and it seems that they will continue to remain as such, since it is unethical to perform a randomized study with a control group of resistant hypertensives that will remain untreated. Since direct data is not available, only clinically meaningful assumptions can
be made based on indirect information and using common sense. Therefore, for the purpose of this paper we'll use data from the past (before the era of antihypertensive therapy), data from clinical studies involving patients with severe or malignant hypertension, data from small clinical studies in patients with resistant hypertension, and from subgroups of patients included in large clinical trials.

This paper attempts to present available evidence regarding the benefits of treatment and control of resistant hypertension, to highlight the significant scarcity of data in this population, and to critically evaluate the use of data from other hypertensive subgroups for extrapolation in resistant hypertension.

2. Data on Malignant Hypertension: Lessons from the Past

The risks of resistant hypertension and the benefits of its management remind one of the story of malignant hypertension. Although uncontrolled or resistant hypertension is a different entity from malignant hypertension, it is well known that in the long term, untreated or uncontrolled hypertension can lead to “accelerated/malignant” phase (VA studies). The term malignant hypertension was introduced by Volhard and Fahr in 1914 for patients with severe hypertension and renal insufficiency [12]. The term was abandoned until the landmark studies performed at Mayo Clinic by Keith and Wagener. It was observed that the prognosis of malignant hypertension was extremely grave. In the first study from Mayo Clinic, only 7 out of 81 patients with malignant hypertension were still alive after the fifty months of followup, while the average length of life was eight months [13]. Retinitis was highlighted as an essential part of malignant hypertension and was significantly associated with mortality; the average length of life in patients with Grade I retinitis was 17 months and that of patients with Grade IV retinitis was 2 months. In a later study of 146 patients, only 1 patient was alive at the end of five-year follow-up period [14]. Further reports of more than 1400 patients with malignant hypertension have confirmed the findings of Mayo Clinic and revealed that the five-year mortality was over 90%, even until the 60s [15–18]. It’s worth noting that no therapy was available at that time. Protein and salt restriction, the rice diet, and mild sedatives were used for the treatment of hypertension; however, the results were all but hopeless regarding long-term improvements [19–23]. Therefore, other therapeutic approaches for malignant hypertension were considered.

Experimental and human studies have revealed the central role of SNS in the pathogenesis of arterial hypertension. Due to the lack of effective therapeutic measures for malignant hypertension, sympathectomy was proposed by many physiologists, such as Pende, Danielopolu, and Jonnesco. Sympathectomy was tested up, that point, for the management of peripheral vascular disease (Jaboulay and Leriche in France), angina pectoris (Jonnesco and Danielopolu in Romania), spastic paralysis (Royle and Hunter in Australia), and Raynaud’s disease and scleroderma in Germany (Bruening). Sympathectomy for the treatment of malignant hypertension appears to have taken place for the first time in Germany as early as 1923 [24]. It was introduced in the US by Alfred Adson at the Mayo Clinic and by Max Peet at Ann Arbor. It was rapidly realized that sympathectomy dramatically increased the survival of patients with malignant hypertension. The five-year mortality rates of patients with Grade IV retinopathy fell from 99% at the Keith Wagener series to 66.5% in sympathectomized patients [25, 26]; similar impressive improvements were observed in patients with Grade III retinopathy (Figure 1).

The pioneer work of Peet, Adson Crile, Hener, Page, Grimson, Hinton, and others was reinforced by Reginald Smithwick, who established the operation worldwide as an effective method of lowering blood pressure in patients with malignant hypertension. Until 1960, a plethora of papers reported the effects of sympathectomy in several thousand patients with malignant hypertension all over the world [27–38], pointing towards dramatic improvements in the survival of operated patients [39] when compared to conservative management (Figure 2).

The indications for sympathectomy waxed and waned during this period. The operation was initially reserved for patients with severe hypertension without significant target organ damage (heart failure, chronic renal disease, angina), was later performed irrespective of the organ damage, and finally restricted to patients without chronic complications since the benefits were more apparent in such patients. Similarly, the extent of the operation varied between the different centers, due to the incomplete understanding of sympathetic anatomy and the absence of appropriate studies comparing the various surgical approaches. The common denominator of all operating techniques was the need for prolonged hospitalization and long recovery period. Another annoying aspect of sympathectomy was the lack of satisfactory predictors of blood pressure response to the operation. Although several tests have been used, the results were inconclusive and sometimes misleading.

The most important limitation of sympathectomy was the safety of the procedure. Adverse events were frequent
and annoying, such as orthostatic hypotension, perioperative pain, orthostatic tachycardia, anhidrosis, intestinal and sexual problems, and palpitations, while more serious complications have been reported, such as perioperative death, stroke, myocardial infarction, paraplegia, and spinal cord injury. The operation was unpleasant and intolerable and many hypertension experts remained skeptical; Ed Weiss stated in 1937 “… and now to cap the climax of his difficulties the unfortunate person with hypertension seems about to fall into the clutches of the neurosurgeon who is prepared to separate him from his sympathetic nervous system”, while Homer Smith used the words “investigation and desperation” for sympathectomy. It was not until the introduction of effective antihypertensive drug therapy that the benefits and risks of sympathectomy were fully reevaluated.

The interest in sympathectomy faded quite suddenly with the advent of antihypertensive therapy. Centrally acting drugs (ganglion-blocking agents, reserpine) have offered similar beneficial effects [40] (Figure 3). The introduction of diuretics has closed the circle of sympathectomy in the treatment of hypertension, highlighting that therapeutic options fade away when new, more promising treatments appear. Of note, blood pressure control significantly affected the survival of treated patients [40] (Figure 4), underlining that uncontrolled hypertension is associated with increased mortality rates.

3. Data from Trials in Severe Hypertension: The VA Study

Despite the impressive benefits of antihypertensive drugs that have established their use in the treatment of malignant hypertension, their role in the treatment of milder forms of hypertension remained controversial for a significant period of time. Even in 1966, it was stated in the book Controversy in Internal Medicine that drug treatment of essential hypertension was not beneficial [41]. Several reasons contributed to the “resistance” of hypertension specialists, primary care physicians, and relevant authorities to recognize the benefits of antihypertensive therapy. Those benefits include the following.

(a) The general belief that vascular changes represent a primary pathologic process that is independent of blood pressure levels. Hypertension was considered to be merely a symptom and not the cause of vascular
complications, therefore the motto “treat the patient, not the manometer” was adopted by the majority.

(b) The inheritance of Sir William Osler promoted therapeutic nihilism. The nihilistic attitude regarding the role of drug therapy may be attributed to Osler’s quote: “one of the first duties of the physician is to educate the masses not to take medicine” [42]. However, this was probably a misinterpretation of Osler’s beliefs, since Osler was referring to the drugs available at his time, the use of which reached the limits of charlatanism, and not modern antihypertensive drugs that were not available at his time.

(c) The special emphasis and the exaggerated focus that were given in secondary forms of hypertension, the prevalence of which was largely overestimated and absorbed the vast majority of available grants. The opinion that one has to find the cause before treating the disease has prevailed, thus rendering “empiric” antihypertensive therapy “a shot in the dark”, an approach that was not appreciated at all. However, the cause of hypertension remained unknown for the vast majority of patients, and it was not unusual for such patients to remain untreated.

(d) The role of preventive medicine was not considered crucial and had not gained wide popularity at that time. Patients, physicians, and the media were not stuck by the benefits of prevention, since the whole society was not ready to move from therapy to prevention.

(e) Maybe the most important factor that restricted the wide adoption of antihypertensive drugs was the lack of convincing clinical studies to verify the benefits of treating essential hypertension.

The first organized data demonstrating benefit from the treatment of severe hypertension came from the Veterans Administration study group. Under the leadership of Edward Freis the first placebo controlled study was carried out in patients with severe hypertension. In that study (published in 1967), 143 patients with severe untreated hypertension (diastolic >115 mmHg) were randomized to either treatment or placebo [43]. In only 20 months, it became apparent that treatment of these patients with severe blood pressure elevation was dramatically beneficial. Twenty-six events occurred in the placebo arm and only 1 in the treated arm (Table 1). It is important to note that 12 out of 26 events were accelerated hypertension leading to malignant hypertension. Since then, the standard of care is to treat severe hypertension; it is unlikely that the study will be repeated. Although the study was placebo controlled, it is reasonable to assume that even treated patients who remain with severe blood pressure elevations (i.e., resistant to treatment) will have similarly bad prognosis.

Confirmation of this assumption comes from many longitudinal studies, cohorts, or subgroup analyses. In the Australian National Blood Pressure study, early in the antihypertensive therapy era, it was shown that patients with uncontrolled blood pressure despite triple therapy had a four-fold increased risk for cardiovascular events compared to patients with controlled blood pressure [44–46].

4. Data from Small Clinical Studies

Virtually no longitudinal study has addressed the particular prognosis of resistant hypertension. Relevant information may be extracted only from small clinical studies. Isakson and Ostergren studied 36 patients with resistant hypertension in Sweden for a 7-year follow-up period [47]. For each of these patients, two control patients were randomly selected from a reference group (retrospectively, matched for age and gender), and the outcomes of the two groups were compared. It has been shown that patients with resistant hypertension had an almost 3-fold increased risk for cardiovascular events (stroke, transient ischemic attacks, myocardial infarction, death, heart failure, renal failure, new onset diabetes) compared to patients with controlled hypertension (odds ratio 2.71; P < .05).

Redon conducted, in Spain, a prospective study of 86 patients with resistant hypertension (diastolic blood pressure >100 mmHg) and a long follow-up period (49 months average) using ambulatory blood pressure measurement (ABPM) [48]. It was reported that patients with poorly controlled blood pressure (daytime diastolic blood pressure >97 mmHg) had more than 6 times higher relative risk for morbid cardiovascular events (relative risk: 6.42; 95% CI: 1.39–29.7; P = .017) compared to patients with relatively controlled blood pressure (daytime diastolic blood pressure <88 mmHg) (Figure 5). It should be noted, however, that the number of patients and events were relatively small, office blood pressure was not independently associated with morbid events, data regarding systolic blood pressure were not provided, and the cut-off limit of daytime diastolic blood pressure (88 mmHg) was higher than what is currently considered normal (85 mmHg).

Pierdomenico in Italy studied a larger number of patients (130 resistant hypertensives) for a slightly longer follow-up period (4.98 ± 2.9 years) using ABPM as well [49]. Moreover, the study compared the outcomes of patients with true resistant hypertension (high clinic and ambulatory blood pressure) to the outcomes of patients with false resistant hypertension (high clinic and normal ambulatory blood pressure).
resistant hypertension according to daytime diastolic blood pressure resistant hypertension using hard endpoints justifies other
trolled versus controlled blood pressure in patients with 
the lack of reliable data regarding the outcome of uncon-
5. Data from Large Clinical Trials

The lack of reliable data regarding the outcome of uncon-
trolled versus controlled blood pressure in patients with 
resistant hypertension using hard endpoints justifies other

approaches. One can use data from other patient populations
and make rational assumptions, although extrapolation

carries inherent risks and has severe limitations.

For example, in a large cohort of hypertensive males
(4,714 patients), it has been shown that cardiovascular
mortality was almost twice as high in male patients
with uncontrolled hypertension compared to patients with well-
controlled blood pressure (risk ratio: 1.66; 95% CI: 1.04–
2.64), although particular data regarding resistant hyper-
tension are not provided [56]. In another cohort of 11,912
veteran male patients followed for 15 years, uncontrolled
hypertension (systolic blood pressure >150 mmHg) was
associated with increased risk of end-stage renal disease
(risk ratio: 3.00; 95% CI: 2.09–4.55; P < .001) [57]. Is this
exaggerated cardiovascular risk of uncontrolled hyperten-
sion applicable in resistant hypertension? Common sense
dictates that there is no reason to assume the opposite.
Until convincing data becomes available, it seems clinically
wise to assume that controlling blood pressure in resistant
hypertension is beneficial, and treating physicians should
make every possible effort towards this direction.

Relevant information can be obtained from large clinical
trials. Although no trial has been specifically designed to
evaluate the benefits of blood pressure control in resistant
hypertension, data from recent large trials regarding patients
that fulfill the definition of resistant hypertension will be
valuable until the conduction of a study devoted to resistant
hypertension. We have to keep in mind, however, the
inherent limitations of such studies, that besides the post-hoc
analysis, they have used unusual antihypertensive regimes,
which are seldom used in everyday clinical practice.

In the ASCOT trial, the combination of older drugs
(diuretics + beta blockers) was compared to newer drugs
(ACEinhibitors + calcium antagonists) [58]. In patients not
achieving blood pressure control, alpha blockers have been
added as third-line and spironolactone as fourth-line ther-
apy. It is obvious that some patients from the diuretic/beta
blocker group may be labeled as resistant hypertensives when
the addition of alpha blockers was ineffective. It should be,
recognized however, that the combination of a diuretic with a
beta- and an alpha-blocker is uncommon in everyday clinical
practice. Calcium antagonists or agents acting on the renin-
angiotensin axis or a combination of both are used for the
vast majority of uncontrolled patients.

Similar problems are applicable to the ACCOMPLISH
trial, which compared the combination of an ACEinhibitor
with diuretics or calcium antagonists [59]. Uncontrolled
patients were allowed to use beta blockers. Therefore patients
from the first group that remained uncontrolled with the
triple combination of ACEinhibitors, diuretics, and beta-
blockers can be characterized as resistant hypertensives
and be used as a source of valuable data extraction. This
combination is more clinically meaningful than the one used
in the ASCOT trial, even this, however, excludes the use of
calcium antagonists, which are among the most commonly
prescribed drugs in the western world for the treatment of
resistant hypertension.

The ALLHAT trial confronts similar problems. Patients
were assigned to receive diuretics, ACEinhibitors, calcium
antagonists, or alpha-blockers, and were allowed to use beta-blockers, clonidine, or hydralazine in case the blood pressure remained above goal [60]. Patients included in the diuretic group that remained uncontrolled despite the use of two additional drugs meet the criteria of resistant hypertension. However, as one can easily notice, the drug combinations that were actually used in this study are seldom used in everyday clinical practice.

6. Conclusions

Data from large clinical trials in different subgroups of hypertensive patients suggest an increased prevalence of resistant hypertension. Data regarding the risks of resistant hypertension, as well as the benefits of treatment and control of blood pressure in resistant hypertensive patients is scarce. However, data from small clinical studies and observational cohorts consistently points towards an increased cardiovascular risk in patients with resistant hypertension. Moreover, available information suggests that there is substantial benefit from appropriate treatment and control of resistant patients. Recent randomized studies in resistant hypertension assessed the efficacy and safety of either new drugs (such as darusentan) [61, 62] or interventional techniques (such as carotid baroreceptor stimulation or renal sympathetic denervation) [63, 64]. We believe that appropriate large, long-term studies are needed to evaluate the prevalence and the risks of resistant hypertension, as well as the significant benefits of treating and controlling resistant hypertension.

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